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# SEARCH REQUEST FORM

## Scientific and Technical Information Center

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Please provide a detailed statement of the Include the elected species or structures, utility of the invention. Define any terms known. Please attach a copy of the cover	e search topic, and describe keywords, synonyms, acro s that may have a special m sheet, pertinent claims, and	as specifically as possible the subject nyms, and registry numbers, and comi eaning. Give examples or relevant cit d abstract.	bine with the concept or tations, authors, etc, if
Title of Invention:	ced transpo	ret using membrane	disruptive agents
Inventors (please provide full names):	Allan J. H	Offman, Patrick	E. S. Stayton,
and Mason My	and 1	Siron Murthy	
Earliest Priority Filing Date://	7/2000	· · · · · · · · · · · · · · · · · · ·	
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Technical Info. Specialist Searcher PhoM/1 6A02 Tel: 308-4491	AA Sequence (#)	Dialog	
Searcher Location:	Structure (#)	Questel/Orbit	
Date Searcher Picked Up: 3.7.02	Bibliographic	Dr.Link	
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Searcher Prep & Review Time:	Fulltext	Sequence Systems	41
Clerical Prep Time:	Patent Family	WWW/Internet	<del></del>
Online Time:	Other	Other (specify)	

PTO-1590 (8-01)

BEST AVAILABLE COPY

Inventor South

Tran 09/755,701

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=> d que 16;d his 17-689 SEA "HOFFMAN A"/AU OR "HOFFMAN A S"/AU 339 SEA "HOFFMAN ALLAN"/AU OR ("HOFFMAN ALLAN S"/AU OR "HOFFMAN L2 ALLAN SACHS"/AU) 218 SEA ("STAYTON P"/AU OR "STAYTON P S"/AU OR "STAYTON PARICK L3 S"/AU OR "STAYTON PAT"/AU OR "STAYTON PAT S"/AU OR "STAYTON PATRICK"/AU OR "STAYTON PATRICK S"/AU OR "STAYTON PATRICK SEAN"/AU) 582 SEA ("MURTHY N"/AU OR "MURTHY N B K"/AU OR "MURTHY N C"/AU OR L4"MURTHY N D A"/AU OR "MURTHY N G K"/AU OR "MURTHY N K"/AU OR "MURTHY N KRISHNA"/AU OR "MURTHY N L"/AU OR "MURTHY N L N"/AU OR "MURTHY N L NARAYANA"/AU OR "MURTHY N M"/AU OR "MURTHY N MANOHARA"/AU OR "MURTHY N N"/AU OR "MURTHY N NARASIMHA"/AU OR "MURTHY N R"/AU OR "MURTHY N RADHA KRISHNA"/AU OR "MURTHY N RAVINDRA"/AU OR "MURTHY N S"/AU OR "MURTHY N S A"/AU OR "MURTHY N S N"/AU OR "MURTHY N S R"/AU OR "MURTHY N S R K"/AU OR "MURTHY N S R KRISHNA"/AU OR "MURTHY N S S"/AU OR "MURTHY N S SATYA"/AU OR "MURTHY N SANJEEVA"/AU OR "MURTHY N SITARAMA"/AU OR "MURTHY N SONJEEVA"/AU OR "MURTHY N SREEDHARA"/AU OR "MURTHY N SRINIVAS"/AU OR "MURTHY N SRINIVASA"/AU OR "MURTHY N SURYANARAYANA"/AU OR "MURTHY N T"/AU OR "MURTHY N V"/AU OR "MURTHY N V A"/AU OR "MURTHY N V ADINARAYANA"/AU OR "MURTHY N V K"/AU OR "MURTHY N V K K"/AU OR "MURTHY N V KISHORE"/AU OR "MURTHY N V KISHORE KUMAR"/AU OR "MURTHY N V S N"/AU OR "MURTHY N V S R"/AU OR "MURTHY N VISHNU"/AU) 21 SEA "MURTHY NIREN"/AU 1753 SEA (L1 OR L2 OR L3 OR L4 OR L5) L5L6

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(FILE 'BIOSIS, HCAPLUS, WPIDS' ENTERED AT 10:22:49 ON 07 MAR 2002)
L7
             55 S L6 AND MEMBRAN?
             58 S TRANSPORT? AND L6
L8
L9
            105 S L7 OR L8
L10
          90005 S DISRUPT?
L11
        1481201 S POLYMER##
L12
             10 S L9 AND L10 AND L11
L13
             13 S L9 AND L10
L14
             37 S L9 AND (L10 OR L11)
L15
             32 DUP; REM L14 (5 DUPLICATES REMOVED)
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FILE 'BIOSIS, HCAPLUS, WPIDS' ENTERED AT 10:27:57 ON 07 MAR 2002

=> d bib ab 1-32

L15 ANSWER 1 OF 32 HCAPLUS COPYRIGHT 2002 ACS DUPLICATE 1 AN 2001:525957 HCAPLUS

```
135:127195
DN
     Enhanced transport of therapeutic and diagnostic agents using
ΤI
     membrane disruptive acid-sensitive polymers
     Hoffman, Allan S.; Stayton, Patrick S.; Murthy,
IN
     University of Washington, USA
PA
     PCT Int. Appl., 50 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
                      KIND
                            DATE
                                          APPLICATION NO. DATE
                            _____
                                           _____
                            20010719
                                           WO 2001-US356
                                                            20010105
PI
     WO 2001051092
                       A2
                            20011206
     WO 2001051092
                      A3
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
         W:
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE; SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,
             ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                            20000107
PRAI US 2000-174893
                      Ρ
     Compns. and methods for transport or release of therapeutic and diagnostic
     agents, metabolites or other analytes from cells, compartments within
     cells, or through cell layers or barriers are described. The compns.
     include a membrane barrier transport enhancing agent and are usually
     administered in combination with an enhancer and/or exposure to stimuli to
     effect disruption or altered permeability, transport or release. In a
     preferred embodiment, the compns. include compds. which disrupt endosomal
     membranes in response to the low pH in the endosomes but which are
     relatively inactive toward cell membranes (at physiol. pH, but can become
     active toward cell membranes if the environment is acidified below pH
     6.8), coupled directly or indirectly to a therapeutic or diagnostic agent.
     Other disruptive agents can also be used, responsive to stimuli and/or
     enhancers other than pH, such as light, elec. stimuli, electromagnetic
     stimuli, ultrasound, temp., or combinations thereof. The compds. can be
     coupled by ionic, covalent or H bonds to an agent to be delivered or to a
     ligand which forms a complex with the agent to be delivered. Agents to be
     delivered can be therapeutic and/or diagnostic agents. Treatments which
     enhance delivery such as ultrasound, iontophoresis, and/or electrophoresis
     can also be used with the disrupting agents. For example, a terpolymer of
     dimethylaminoethyl methacrylate, Bu methacrylate, and styrene benzaldehyde
     was prepd. for the membrane-disruptive backbone which was then PEGylated
     with thiol-terminated monofunctional or heterofunctional PEGs. The
     acid-degradable linkage was a p-aminobenzaldehyde acetal.
    ANSWER 2 OF 32 WPIDS COPYRIGHT 2002
T.15
                                            DERWENT INFORMATION LTD
     2001-441402 [47]
AN
                       WPIDS
    C2001-133267
DNC
TΤ
     Drug delivery system for controlled release of active agent in
     gastro-intestinal tract comprises a matrix consisting of (non)degradable
     polymer, (non) continuous membrane and drug.
DC
     A96 B07
     FRIEDMAN, M; HOFFMAN, A; KLAUSNER, E; LAVY, E
IN
PΑ
    (YISS) YISSUM RES DEV CO HEBREW UNIV JERUSALEM
CYC
     WO 2001037812 A2 20010531 (200147)* EN
                                              46p
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RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR L\$ LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001016477 A 20010604 (200153)

ADT WO 2001037812 A2 WO 2000-IL774 20001120; AU 2001016477 A AU 2001-16477 20001120

FDT AU 2001016477 A Based on WO 200137812

PRAI IL 1999-133196 19991129

AB WO 200137812 A UPAB: 20010822

NOVELTY - A gastro-retentive drug delivery system for the controlled release of an active agent in the gastrointestinal (GI) tract comprises a single or multi-layered matrix having a 2D or 3D configuration comprising a polymer that retains in the stomach more than a conventional dosage form, a continuous or non-continuous membrane and a drug that may be particulate or contained in a drug-containing form.

DETAILED DESCRIPTION - A pharmaceutical gastro-retentive drug delivery system for the controlled release of an active agent in the GI tract comprises a single or multi-layered matrix having a 2D or 3D configuration comprising a polymer that does retain in the stomach more than a conventional dosage form, a continuous or non-continuous membrane and a drug that may be particulate or contained in a drug-containing form. The polymer of the matrix is a degradable polymer of a hydrophilic polymer that is not readily soluble in gastric fluids, an enteric polymer substantially insoluble at pH less than 5.5 and/or a hydrophobic polymer, and/or a degradable polymer. The membrane does not retain in the stomach more than a conventional

membrane does not retain in the stomach more than a conventional dosage form, affixed or attached to the matrix, and is a **polymer** with substantial mechanical strength. The drug component is embedded or entrapped in the matrix or is attached to the delivery system and remains in the stomach for 3-24 hours.

An INDEPENDENT CLAIM is also included for the delivery system in the form of a capsule.

USE - A drug release system for the controlled release of an active agent in the GI tract.

ADVANTAGE - Improved efficiency of treatment by reducing the frequency of administration and the application of single doses for improved patient compliance.

DESCRIPTION OF DRAWING(S) - The controlled release drug delivery system.

Controlled release drug delivery system 1

3D matrix containing the drug 100

Strips (fixed to the sides of the matrix to form a continuous  ${\tt membrane})\ 110$  .

Shielding layer 120

Anti-adhering powder layer 130

Dwg.1/5

- L15 ANSWER 3 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 2001:355468 HCAPLUS
- DN 135:2504
- TI Size-dependent control of the binding of biotinylated proteins to streptavidin using a **polymer** shield
- AU Ding, Zhongil; Fong, Robin B.; Long, Cynthia J.; Stayton, Patrick S.; Hoffman, Allan S.
- CS Department of Bioengineenng, University of Washington, Seattle, WA, 98195, USA

- SO Nature (London, U. K.) (2001), 411(6833), 59-62 CODEN: NATUAS; ISSN: 0028-0836
- PB Nature Publishing Group
- DT Journal LA English
- Many medical and biotechnol. processes rely on controlling and AB manipulating the mol.-recognition capabilities of proteins. This can be achieved using small mols. capable of competing for protein binding or by changing environmental parameters that affect protein structure and hence binding. An alternative is provided by stimuli-responsive polymers that change reversibly from a water-sol. expanded coil to a water-insol. collapsed globule upon small changes in temp., pH or light intensity: when attached to proteins in the vicinity of their binding sites, they reversibly block and release small ligands. Here we show how this approach can be extended to achieve size-selective binding of large, macromol. ligands. We use the thermally responsive polymer poly(N,N-diethylacrylamide) (PDEAAm), and attach it to the protein streptavidin approx. 20 .ANG. from the binding site for biotinylated proteins. Below the lower crit. soln. temp. of PDEAAm, the polymer is in its extended state and acts as a 'shield' to block the binding of large biotinylated proteins; above this temp., it collapses and exposes the binding site, thereby allowing binding. We find that the degree of shielding depends on both the size of the biotinylated protein and the size of PDEAAm, suggesting that 'smart' polymer shields could be tailored to achieve a wide range of size-dependent ligand discrimination for use in affinity sepns., biosensors and diagnostics technologies.
- RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L15 ANSWER 4 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 2001:406498 BIOSIS
- DN PREV200100406498
- TI Control of shape and size of vascular smooth muscle cells in vitro by plasma lithography.
- AU Goessl, Andreas; Bowen-Pope, Daniel F.; Hoffman, Allan S. (1)
- CS (1) Department of Bioengineering, University of Washington, Seattle, WA, 98195: hoffman@u.washington.edu USA
- SO Journal of Biomedical Materials Research, (October, 2001) Vol. 57, No. 1, pp. 15-24. print.
  ISSN: 0021-9304.
- DT Article
- LA English
- SL English
- The ability to control the shape and size of cells is an important AR enabling technique for investigating influences of geometrical variables on cell physiology. Herein we present a micropatterning technique ("plasma lithography") that uses photolithography and plasma thin-film polymerization for the fabrication of cell culture substrates with a cell-adhesive pattern on a cell-repellent (non-fouling) background. The micron-level pattern was designed to isolate individual vascular smooth muscle cells (SMC) on areas with a projected area of between 25 and 3600 mum2 in order to later study their response to cytokine stimulation in dependence of the cell size and shape as an indication for the phenotypic state of the cells. Polyethylene terephthalate substrates were first coated with a non-fouling plasma polymer of tetraglyme (tetraethylene glycol dimethyl ether). In an organic lift-off process, we then fashioned square- and rectangular-shaped islands of a thin fluorocarbon plasma polymer film of apprx12-nm thickness. Electron spectroscopy for chemical analysis and secondary ion mass spectroscopy were used to optimize the deposition conditions and

characterize the resulting polymers. Secondary ion mass spectroscopy imaging was used to visualized the spatial distribution of the polymer components of the micropatterned surfaces. Rat vascular SMC were seeded onto the patterned substrates in serum-free medium to show that the substrates display the desired properties, and that cell shape can indeed be controlled. For long-term maintenance of these cells, the medium was augmented with 10% calf serum after 24 h in culture, and the medium was exchanged every 3 days. After 2 weeks, the cells were still confined to the areas of the adhesive pattern, and when one or more cells spanned more than one island, they did not attach to the intervening tetraethylene glycol dimethyl ether (tetraglyme) background. Spreading-restricted cells formed a well-ordered actin skeleton, which was most dense along the perimeter of the cells. The shape of the nucleus was also influenced by the pattern geometry. These properties make the patterned substrates suitable for investigating if the phenotypic reversion of SMC can be influenced by controlling the shape and size of SMC in vitro.

- L15 ANSWER 5 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 2000:594116 HCAPLUS
- DN 133:325550
- TI Bioinspired polymeric conjugates for biotechnologies
- AU Stayton, Patrick S.; Hoffman, Allan S.; Murthy, Niren; Cheung, Charles; Lackey, Chantal; Ding, Zhongli; Shimoboji, Tsuyoshi; Press, Oliver
- CS Department of Bioengineering, University of Washington, Seattle, WA, 98195, USA
- SO Polym. Prepr. (Am. Chem. Soc., Div. Polym. Chem.) (2000), 41(2), 1607-1608 CODEN: ACPPAY; ISSN: 0032-3934
- PB American Chemical Society, Division of Polymer Chemistry
- DT Journal
- LA English
- AB The authors developed polymeric systems to manipulate intracellular trafficking by enhancing transport across the endosomal membrane. Bioinspired polymeric carriers for cytoplasmic delivery of genes and proteins were designed.
- RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L15 ANSWER 6 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 2001:397371 HCAPLUS
- DN 136:147277
- TI Measured bioeffects of tone-burst ultrasound in combination with poly(propyl acrylic) acid (PPAA)
- AU Porter, Tyrone; Hadley, Maile; Nickerson, Josh; Mourad, Pierre; Crum, Lawrence; Murthy, Niren; Stayton, Patrick; Hoffman, Allan
- CS Applied Physics Laboratory, Seattle, WA, 98105, USA
- SO Proceedings IEEE Ultrasonics Symposium (2000), (Vol. 2), 1359-1362 CODEN: PIEUEZ; ISSN: 1051-0117
- PB Institute of Electrical and Electronics Engineers
- DT Journal
- LA English
- AB In this study, High Intensity Focused Ultrasound (HIFU) is combined with the pH-sensitive cell membrane disrupting polymer PPAA (poly-Pr acrylic acid) at sublethal doses to achieve hemolysis of human erythrocytes and sonoporation of suspended cells. For our studies, a 1 mL sample of cells suspended in phosphate buffered saline (PBS) was simultaneously exposed to 1.1 MHz acoustic tone bursts and PPAA at a temp. of 37.degree.. We vary the pH of the suspension fluid and amt. of PPAA added to assess the

influence of its structural conformation, functionality, and concn. upon measured bioeffects. For hemolysis study, we suspended erythrocytes at a final concn. of 108 cells/mL. Damage to the cell suspension was detd. by measuring the amt. of Hb released using a spectrophotometer. A passive cavitation detection system was utilized to monitor the acoustic emissions from the cell suspension during exposure to ultrasound. In the presence of PPAA, there is a significant increase in cavitation and bioeffects during ultrasound exposure at more acidic pH levels. This polymer/ultrasound synergy is pH independent, unlike the synergy of ultrasound with poly(Et acrylic acid). The levels of cavitation and hemolysis measured from HIFU/PPAA synergy was compared with levels measured from HIFU/Optison synergy to assess the effectiveness of the polymer in an acoustic field.

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L15 ANSWER 7 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 2000:208542 HCAPLUS
- DN 133:109744
- TI pH sensitive membrane disruptive PEGylated polycations
- AU Murthy, Niren, Stayton, Patrick S.; Hoffman, Allan S.
- CS Department of Bioengineering, University of Washington, Seattle, WA, 98195, USA
- SO Polym. Prepr. (Am. Chem. Soc., Div. Polym. Chem.) (2000), 41(1), 1010-1011 CODEN: ACPPAY; ISSN: 0032-3934
- PB American Chemical Society, Division of Polymer Chemistry
- DT Journal
- LA English
- AB A new method for the synthesis of novel PEGylated pH sensitive membrane-disruptive polycations as potential oligonucleotide delivery vehicles has been presented. The strategy is based on grafting PEG onto a hydrophobic-polycationic backbone through an acid degradable acetal linkage. The acetal linkage used for the PEGylation of Copolymer I had a half life of 15 min at pH 5.4, but at pH 7.4 less than 10% of the acetals were hydrolyzed after 80 min. Copolymer I has a hydrolysis rate suitable for drug delivery purposes. The hydrolysis of the PEG grafts and activation of its membrane disruptive activity occur in less than 20 min at pH 5.0. Copolymer I was membrane disruptive at pH 5.0 but not at pH 7.4. The above copolymers should therefore have applications for the delivery of neg. charged polyanions such as DNA or ODNs to cells.
- RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L15 ANSWER 8 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 2000:334740 HCAPLUS
- TI pH-Sensitive membrane disruptive PEGylated polycations.
- AU Murthy, Niren; Stayton, Patrick; Hoffman, Allan
- CS Bioengineering; University of Washington, Seattle, WA, 98195, USA
- SO Book of Abstracts, 219th ACS National Meeting, San Francisco, CA, March 26-30, 2000 (2000), POLY-551 Publisher: American Chemical Society, Washington, D. C. CODEN: 69CLAC
- DT Conference; Meeting Abstract
- LA English
- AB Oligonucleotides (ODNs) complementary to specific mRNAs have shown considerable promise for the treatment of diseases such as cancer and viral infections. However, the clin. use of ODNs has been hindered by the

lack of an effective ODN delivery vehicle. PEGylated polycations have recently been considered as an ODN delivery vehicle. PEGylated polycations form polyelectrolyte complexes (PECs) with ODNs. However, the efficiency of pegylated PECs in delivering ODNs is not optimal because endocytosed pegylated PECs are trafficked and degraded in lysosomes. In this report a movel method is developed for the synthesis of PEGylated hydrophobic polycations which have the PEG groups conjugated to the backbone polycations via acid degradable acetal bonds. Thus the pegylated polymers should lose their PEGs and become hydrophobic and membrane disruptive at the endosomal pHs of 6.0 and below, but should not be membrane disruptive at the plasma pH of 7.4. These novel polymers should be useful for delivery of endocytosed anionic drugs such as ODNs and DNA.

- L15 ANSWER 9 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 2000:191375 BIOSIS
- DN PREV200000191375
- Molecular engineering of proteins and polymers for targeting and intracellular delivery of therapeutics.
- AU Stayton, Patrick S. (1); Hoffman, Allan S.;
  Murthy, Niren; Lackey, Chantal; Cheung, Charles; Tan, Philip;
  Klumb, Lisa A.; Chilkoti, Ashutosh; Wilbur, F. Scott; Press, Oliver W.
- CS (1) Department of Bioengineering, University of Washington, Seattle, WA, 98195 USA
- SO Journal of Controlled Release, (March 1, 2000) Vol. 65, No. 1-2, pp. 203-220.
  ISSN: 0168-3659.
- DT General Review
- LA English
- SL English
- AB There are many protein and DNA based therapeutics under development in the biotechnology and pharmaceutical industries. Key delivery challenges remain before many of these biomolecular therapeutics reach the clinic. Two important barriers are the effective targeting of drugs to specific tissues and cells and the subsequent intracellular delivery to appropriate cellular compartments. In this review, we summarize protein engineering work aimed at improving the stability and refolding efficiency of antibody fragments used in targeting, and at constructing new streptavidin variants which may offer improved performance in pre-targeting delivery strategies. In addition, we review recent work with pH-responsive polymers that mimic the membrane disruptive properties of viruses and toxins. These polymers could serve as alternatives to fusogenic peptides in gene therapy formulations and to enhance the intracellular delivery of protein therapeutics that function in the cytoplasm.
- L15 ANSWER 10 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 2000:533806 BIOSIS
- DN PREV200000533806
- TI Characterization and gene therapy applications of pH-responsive, membrane-active polymers.
- AU Black, F. E. (1); Cheung, C. Y. (1); Murthy, N. (1); Hoffman, A. S. (1); Stayton, P. S. (1)
- CS (1) Department of Bioengineering, University of Washington, Seattle, WA, 98195 USA
- SO Journal of Pharmacy and Pharmacology, (September, 2000) Vol. 52, No. Supplement, pp. 42. print.

  Meeting Info.: 137th British Pharmaceutical Conference Birmingham, England, UK September 10-13, 2000
  ISSN: 0022-3573.
- DT Conference

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LA
     English
SL
     English
     ANSWER 11 OF 32 HCAPLUS COPYRIGHT 2002 ACS
L15
                                                       DUPLICATE 2
AN
     1999:451212 HCAPLUS
DN
     131:106813
TI
     Enhanced transport using membrane disruptive
     agents
     Hoffman, Allan S.; Stayton, Patrick; Press, Oliver;
IN
     Tirrell, David; Murthy, Niren; Lackey, Chantal; Crum, Lawrence
     A.; Mourad, Pierre D.; Porter, Tyrone M.
PA
     University of Washington, USA; University of Massachusetts
     PCT Int. Appl., 54 pp.
SO
     CODEN: PIXXD2
DΤ
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
                     KIND
                            DATE
                                           APPLICATION NO.
                                                            DATE
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     WO 9934831
                       A1
                            19990715
                                           WO 1999-US122
                                                            19990105
         W: AU, CA,
                    JΡ
         RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
     AU 9920261
                       A1
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                                                            19990105
     EP 1044021
                       A1
                            20001018
                                           EP 1999-900750
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                            19990105
     Compns. and methods for transport or release of therapeutic and diagnostic
AΒ
     agents or metabolites or other analytes from cells, compartments within
     cells, or through cell layers or barriers are described. The compns.
     include a membrane barrier transport enhancing agent and are usually
     administered in combination with an enhancer and/or exposure to stimuli to
     effect disruption or altered permeability, transport or release.
     preferred embodiment, the compns. include compds. which disrupt endosomal
    membranes in response to the low pH in the endosomes but which are
    relatively inactive toward cell membranes, coupled directly or indirectly
    to a therapeutic or diagnostic agent. Other disruptive agents can also be
    used, responsive to stimuli and/or enhancers other than pH, such as light,
    elec. stimuli, electromagnetic stimuli, ultrasound, temp., or combinations
    thereof.
               The compds. can be coupled by ionic, covalent or H bonds to an
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conjugate gave 70% lysis at 100 .mu.g. RE.CNT 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

agent to be delivered or to a ligand which forms a complex with the agent

diagnostic agents. Treatments which enhance delivery such as ultrasound, iontophoresis, and/or electrophoresis can also be used with the disrupting

The ability of the GALA peptide to lyse erythrocytes was compared

to be delivered. Agents to be delivered can be therapeutic and/or

with that of an GALA poly (acrylic acid) conjugate at pH 5.0. The

L15 ANSWER 12 OF 32 HCAPLUS COPYRIGHT 2002 ACS

AN 1999:780354 HCAPLUS

DN 132:10527

TI Interactive molecular conjugates

IN Hoffman, Allan S.; Stayton, Patrick S.

A University of Washington, USA

SO U.S., 32 pp.

CODEN: USXXAM

DT Patent LA English

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI US 5998588 A 19991207 US 1996-697904 19960830

- AB The combination of the capabilities of stimuli-responsive components such as polymers and interactive mols. to form site-specific conjugates which are useful in a variety of assays, sepns., processing, and other uses is disclosed. The polymer chain conformation and vol. can be manipulated through alteration in pH, temp., light, or other stimuli. The interactive mols. can be biomols. like proteins or peptides, such as antibodies, receptors, or enzymes, polysaccharides or glycoproteins which specifically bind to ligands, or nucleic acids such as antisense, ribozymes, and aptamers, or ligands for org. or inorg. mols. in the environment or manufg. processes. The stimuli-responsive polymers are coupled to the recognition biomols. at a specific site so that the polymer can be manipulated by stimulation to alter ligand-biomol. binding at an adjacent binding site, for example, the biotin binding site of streptavidin, the antigen-binding site of an antibody or the active, substrate-binding site of an enzyme. Binding may be completely blocked (i.e., the conjugate acts as an on-off switch) or partially blocked (i.e., the conjugate acts as a rheostat to partially block binding or to block binding only of larger ligands). Once a ligand is bound, it may also be ejected from the binding site by stimulating one (or more) conjugated polymers to cause ejection of the ligand and whatever is attached to it. Alternatively, selective partitioning, phase sepn. or pptn. of the polymer-conjugated biomol. can be achieved through exposure of the stimulus-responsive component to an appropriate environmental stimulus.
- RE.CNT 96 THERE ARE 96 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L15 ANSWER 13 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1999:246745 BIOSIS
- DN PREV199900246745
- TI Mice that lack the angiogenesis inhibitor, thrombospondin 2, mount an altered foreign body reaction characterized by increased vascularity.
- AU Kyriakides, Thémis R.; Leach, Kathleen J.; Hoffman, Allan S.; Ratner, Buddy D.; Bornstein, Paul (1)
- CS (1) Department of Biochemistry, University of Washington, Seattle, WA, 98195 USA
- SO Proceedings of the National Academy of Sciences of the United States of America, (April 13, 1999) Vol. 96, No. 8, pp. 4449-4454.
  ISSN: 0027-8424.
- DT Article
- LA English
- SL English
- Disruption of the thrombospondin 2 gene (Thbs2) in mice results in a complex phenotype characterized chiefly by abnormalities in fibroblasts, connective tissues, and blood vessels. Consideration of this phenotype suggested to us that the foreign body reaction (FBR) might be altered in thrombospondin 2 (TSP2)-null mice. To investigate the participation of TSP2 in the FBR, polydimethylsiloxane (PDMS) and oxidized PDMS (ox-PDMS) disks were implanted in TSP2-null and control mice. Growth of TSP2-null and control skin fibroblasts in vitro also was evaluated on both types of disks. Normal fibroblasts grew as a monolayer on both surfaces, but attachment of the cells to ox-PDMS was weak and sensitive to movement. TSP2-null fibroblasts grew as aggregates on both surfaces, and

their attachment was further compromised on ox-PDMS. After a 4-week implantation period, both types of PDMS elicited a similar FBR with a collagenous capsule in both TSP2-null and control mice. However, strikingly, the collagenous capsule that formed in TSP2-null mice was highly vascularized and thicker than that formed in normal mice. In addition, abnormally shaped collagen fibers were observed in capsules from mutant mice. These observations indicate that the presence or absence of an extracellular matrix component, TSP2, can influence the nature of the FBR, in particular its vascularity. The expression of TSP2 therefore could represent a molecular target for local inhibitory measures when vascularization of the tissue surrounding an implanted device is desired.

- L15 ANSWER 14 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1999:281611 BIOSIS
- DN PREV199900281611
- TI Hemolytic activity of pH-responsive polymer-streptavidin bioconjugates.
- AU Lackey, Chantal A.; Murthy, Niren; Press, Oliver W.; Tirrell, David A.; Hoffman, Allan S. (1); Stayton, Patrick S.
- CS (1) Department of Bioengineering, University of Washington, Seattle, WA,
- SO Bioconjugate Chemistry, (May-June, 1999) Vol. 10, No. 3, pp. 401-405. ISSN: 1043-1802.
- DT Article
- LA English
- SL English
- Drug delivery systems that increase the rate and/or quantity of drug AΒ release to the cytoplasm are needed to enhance cytosolic delivery and to circumvent nonproductive cell trafficking routes. We have previously demonstrated that poly(2-ethylacrylic acid) (PEAAc) has pH-dependent hemolytic properties, and more recently, we have found that poly(2-propylacrylic acid) (PPAAc) displays even greater pH-responsive hemolytic activity than PEAAc at the acidic pHs of the early endosome. Thus, these polymers could potentially serve as endosomal releasing agents in immunotoxin therapies. In this paper, we have investigated whether the pH-dependent membrane disruptive activity of PPAAc is retained after binding to a protein. We did this by measuring the hemolytic activity of PPAAc-streptavidin model complexes with different protein to polymer stoichiometries. Biotin was conjugated to amine-terminated PPAAc, which was subsequently bound to streptavidin by biotin complexation. The ability of these samples to disrupt red blood cell membranes was investigated for a range of polymer concentrations, a range of pH values, and two polymer -to-streptavidin ratios of 3:1 and 1:1. The results demonstrate that (a) the PPAAc-streptavidin complex retains the ability to lyse the RBC lipid bilayers at low pHs, such as those existing in endosomes, and (b) the hemolytic ability of the PPAAc-streptavidin complex is similar to that of the free PPAAc:
- L15 ANSWER 15 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.DUPLICATE
- AN 2000:9745 BIOSIS
- DN PREV200000009745
- TI The design and synthesis of polymers for eukaryotic membrane disruption.
- AU Murthy, Niren; Robichaud, John R.; Tirrell, David A.; Stayton, Patrick S.; Hoffman, Allan S. (1)
- CS (1) Department of Bioengineering, University of Washington, Seattle, WA, 98195 USA

- Journal of Controlled Release, (Aug. 27, 1999) Vol. 61, No. 1-2, pp. SO 137-143. ISSN: 0168-3659.
- DTArticle
- LA English
- English SL
- The intracellular trafficking of drugs is critical to the efficacy of AB drugs that are susceptible to attack by lysosomal enzymes. It is therefore an important goal to design and synthesize molecules which can enhance the transport of endocytosed drugs from the endosomal compartments to the cytoplasm. The pH of an endosome is lower than that of the cytosol by one to two pH units, depending on the stage of endosomal development. This pH gradient is a key factor in the design of membranedisruptive polymers which could enhance the endosomal release of drugs. Such polymers should disrupt lipid bilayer membranes at pH 6.5 and below, but should be non-lytic at pH 7.4. We have designed and synthesized pH-sensitive synthetic polymers which efficiently disrupt red blood cells within a sharply defined pH range. One of these polymers, poly(ethyl acrylic acid) (PEAAc) has been previously shown to disrupt synthetic vesicles in a pH-dependent fashion (6). PEAAc hemolyzes red blood cells with an activity of 107 molecules per red blood cell, which is as efficient on a molar basis as the peptide melittin. The mechanism of RBC hemolysis by PEAAc is consistent with the colloid osmotic mechanism. PEAAc's hemolytic activity rises rapidly as the pH decreases from 6.3 to 5.0, and there is no hemolytic activity at pH 7.4. A related polymer, poly(propyl acrylic acid) (PPAAc), was synthesized to test whether making the pendant alkyl group more hydrophobic by adding one methylene group would increase the hemolytic activity. PPAAc was found to disrupt red blood cells 15 times more efficiently than PEAAc at pH 6.1. PPAAc was also not active at pH 7.4 and displayed a pH-dependent hemolysis that was shifted toward higher pH's. Random 1:1 copolymers of ethyl acrylate (EA) and acrylic acid (AAc) (which contain random -COOH and -C2H5 groups that are present and regularly repeat in PEAAc) also displayed significant hemolytic activity, with an efficiency close to PEAAc. These results demonstrate that pH-sensitive synthetic polymers can be molecularly engineered to efficiently disrupt eukaryotic membranes within defined and narrow pH ranges. Thus, these polymers might serve as endosomal disruptive agents with specificities for early or late endosomes.
- ANSWER 16 OF 32 HCAPLUS COPYRIGHT 2002 ACS L15
- 1998:481771 <u>HCAPLUS</u> AN
- DN 129:221141
- Design of polymers to increase the efficiency of endosomal TΙ release of drugs
- Murthy, N.; Robichaud, J.; Stayton, P. S.; Press, O. ΑU W.; Hoffman, A. S.; Tirrell, D. A.
- CS Departments of Bioengineering and Medicine, University of Washington, Seattle, WA, USA
- Proc. Int. Symp. Controlled Release Bioact. Mater. (1998), 25th, 224-225 SO CODEN: PCRMEY; ISSN: 1022-0178
- PBControlled Release Society, Inc.

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- DTJournal
- LAEnglish
- The authors prepd. and investigated the ability of 4 pH-sensitive acrylic AΒ polymers to disrupt lipid bilayer membranes by measuring their ability to hemolyze red blood cells. The potential drug delivery polymers were poly(ethacrylic acid), poly(2-propylacrylic acid). poly(2-butylacrylic acid), and acrylic acid-ethacrylic acid copolymer.

- L15 ANSWER 17 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- 1998:481705 HCAPLUS AN
- 129:235484 DN
- Stimuli-responsive biomolecular conjugates: controlled membrane TΙ disruption
- Lackey, C. A.; Murthy, N.; Stayton, P. S.; Press, O. ΑU W.; Hoffman, A. S.; Tirrell, D. A.
- Departments of Broengineering and Medicine, University of Washington, CS Seattle, WA, USA
- SO Proc. Int. Symp. Controlled Release Bioact. Mater. (1998), 25th, 87-88 CODEN: PCRMEY; ISSN: 1022-0178
- Controlled Release Society, Inc. 'PB
- DTJournal
- LA English
- Poly(ethylacrylic acid) complexed to a protein retains its ability to lyse AΒ lipid bilayers at low pH's, such as those existing in endosomes. demonstrates potential to improve proper therapeutic delivery by facilitating endosomal release.
- ANSWER 18 OF 32 HCAPLUS COPYRIGHT 2002 ACS L15
- 1997:274469 HCAPLUS AN
- 126:317781 DN
- Hybrid biomaterials prepared by ozone-induced polymerization. I. Ozonation ΤI of microporous polypropylene
- ΑU Gatenholm, P.; Ashida, T.; Hoffman, A. S.
- Dep. Polymer Technology, Chalmers Univ. Technology, Goteborg, S-412 96, CS
- J. Polym. Scr., Part A: Polym. Chem. (1997), 35(8), 1461-1467 CODEN: JPACEC, ISSN: 0887-624X SO
- PB Wiley
- Journal DT
- LA English
- Exposure of isotactic polypropylene (PP) to ozone resulted in surface AR oxidn., as detected by ESCA, and the formation of peroxides and hydroperoxides. The amt. of oxygen-bearing moieties, as detected by FT-IR, was increased when a microporous membrane with a large surface area was used. Ozonation for an extended period of time, 1-2 h, resulted in a degrdn. of microporous PP, obsd. with SEM as an enlargement of pores and brittle characteristics of the material. The mol. wt. of PP was dramatically reduced after as little as 5 min of ozonation. Exposure to ozone for longer periods of time contributed to further redns. of the mol. wt. and gradual modification of the chem. compn. of PP, restricted, however, to the surface or intercryst. amorphous regions. It was possible to graft 2-hydroxyethyl methacrylate to the ozonated samples, such that the graft copolymer acted as a continuous matrix consequently linked to and reinforced by the PP crystals.
- ANSWER 19 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15
- 1995:293762 BİOSIS ΑN
- DN PREV199598308062
- ΤI Platelet and monoclonal antibody binding to fibrinogen adsorbed on glow-discharge tdeposited polymers.
- ΑU Kiaei, David; Hoffman, Allan S. (1); Horbett, Thomas A.; Lew,
- CS (1) Cent. Bioengineering, FL-20, University Washington, Seattle, WA 98195
- SO Journal of Biomedical Materials Research, (1995) Vol. 29, No. 6, pp. 729-739.
  - ISSN: 0021-9304.

- DT Article
- LA English
- The state of fibrinogen adsorbed on untreated and glow-discharge-treated AB surfaces was examined by measuring platelet adhesion, monoclonal antibody (mAb) binding, the amount of fibrinogen adsorbed, and the amount of adsorbed fibringen which could be eluted with sodium dodecyl sulfate (SDS). Tetrafluoroethylene (TFE) glow-discharge-treated polymers have a lower surface free energy (in air) and retain a larger fraction of adsorbed fibrinogen than untreated surfaces after SDS elution. Platelet adhesion was lowest on the TFE-treated surfaces which retain the highest amounts of fibrinogen after SDS elution. Fibrinogen may undergo unfolding or spreading on the TFE-treated surfaces to minimize interfacial free energy (in water) and maximize protein-surface interactions. When it is adsorbed on the TFE-treated surfaces, fibrinogen evidently assumes a state which somehow prevents its recognition and binding by platelet receptors. Monoclonal antibodies that bind to the three regions in fibrinogen thought to be involved in platelet adhesion were therefore used to detect changes in adsorbed fibrinogen. These regions and the antibodies which bind to them are: the COOH-terminal of the gamma-chain, mAb Ml; the RGD peptide sequence at A-alpha 95-98, mAb R1; the RGD sequence at Aa 572-575, mAb R2. For fibrinogen adsorbed on the untreated or TFE-treated surfaces, Ml and R2 binding was relatively high compared to background, while R1 binding was low. However, the amount of binding of each mAb to fibrinogen adsorbed on the TFE-treated surfaces was equal to or greater than fibrinogen adsorbed to the untreated surfaces. Therefore, antibody-detectable changes in the platelet binding regions of adsorbed fibrinogen that might have been caused by conformational or orientational rearrangements were not observed for the TFE-treated surfaces. The data suggest that the tight binding of fibrinogen on a surface may directly affect the ability of the fibrinogen to interact with the platelet receptors-i.e., that fibrinogen must be loosely held to facilitate maximal interaction with platelet receptors.
- L15 ANSWER 20 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1995:403291 BÍOSIS
- DN PREV199598417591
- TI Silicone-based microcarriers: Preparation and BHK cell culture.
- AU Denkbas, Emir B.; Hoffman, Allan S.; Piskin, Erhan (1)

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- CS (1) Hacettepe Univ., Chem. Engineering Dep., Bioengineering Div., Ankara Turkey
- SO Chemical Engineering Journal, (1995) Vol. 58, No. 1, pp. 65-70. ISSN: 0923-0467.
- DT Article
- LA English
- L15 ANSWER 21 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. DUPLICATE
- AN 1994:271759 BIOSIS
- DN PREV199497284759
- TI Activated, N-substituted acrylamide polymers for antibody coupling: Application to a novel membrane-based immunoassay.
- AU Monji, Nobuo (1); Cole, Carol-Ann; Hoffman, Allan S.
- CS (1) Genet. Systems Corp., 6565 185th Avenue NE, Redmond, WA 98052 USA
- SO Journal of Biomaterials Science Polymer Edition, (1994) Vol. 5, No. 5, pp. 407-420.
  ISSN: 0920-5063.
- DT Article
- LA English
- AB A room-temperature-precipitable, activated terpolymer consisting of N-isopropylacrylamide (NIPAAm)/N-n-butylacrylamide(nBAAm)/N-

acryloxysuccinimide(NASI) (LCST = 7-13 degree C) at a monomer feed ratio of 60:40:2.5, respectively, was prepared and conjugated to an antibody. The conjugate was evaluated in a novel cellulose acetate (CA) membrane-based immunoassay which utilizes the especially strong physical attachment of the polymer to CA to bind and concentrate the polymer attached protein onto the membrane. When compared in the CA membrane immunoassay to the antibody-poly(NIPAAm) conjugate prepared via anhydrous copolymerization of NIPAAm and NASI at the monomer feed ratio of 40: 1, respectively, the performance of the NIPAAm/nBAAm/NASI terpolymer was superior to that of the NIPAAm/NAST copolymer (LCST = 32 degree C) when the studies were carried out at room temperature. However, the terpolymer and copolymer gave equivalent performance when the assay mixture was heated to 45 degree C. These results indicate the importance of the LCST of the polymer component of the Ab-polymer conjugate to its adsorption and binding on the CA membrane.

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L15 ANSWER 22 OF 32 HCAPLUS COPYRIGHT 2002 ACS
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AN 1993:651570 HCAPLUS

DN 119:251570

- TI Novel biomaterials prepared by ozone-induced polymerization
- AU Gatenholm, P. Ashida, T.; Nabeshima, Y.; Hoffman, A. S.
- CS Cent. Bioeng. Univ. Washington, Seattle, WA, 98195, USA
- SO Polym. Mater. Sci. Eng. (1992), 66, 445-6 CODEN: PMSEDG; ISSN: 0743-0515
- DT Journal
- LA English
- AB The O3-induced graft polymn. of hydroxyethyl methacrylate or N-isopropyladrylamide onto the polypropylene films and microporous membranes was carried out to prep. membrane hydrogel hybrid materials for the immobilization of biolog. active species.
- L15 ANSWER 23 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1993:70604 BIOSIS
- DN PREV199395035104
- TI Tight binding of albumin to glow discharge treated polymers.
- AU Kiaei, David; Hoffman, Allan S.; Horbett, Thomas A.
- CS Cent. Bioengineering, Dep. Chemical Engineering, Univ. Washington, Seattle, Washington 98195 USA
- SO Journal of Biomaterials Science Polymer Edition, (1992) Vol. 4, No. 1, pp. 35-44.
  ISSN: 0920-5063.
- DT Article
- LA English
- Tetrafluoroethylene (TFE) glow discharge-treated Dacron vascular grafts resist thrombus deposition, embolization and thrombotic occlusion. In addition, albumin adsorbed on TFE-treated surfaces resists elution by sodium dodecyl sulfate (SDS). Since the tight binding of albumin to TFE-treated surfaces may contribute to their thromboresistant character, we decided to examine the mechanism responsible for this tenacious adsorption. We have investigated albumin adsorption and retention (after SDS elution) on a number of untreated and glow discharge-treated surfaces. Fluorocarbon glow discharge-treated polymers retain a larger fraction of the adsorbed albumin than ethylene and hexamethyldisiloxane glow discharge treated surfaces. Albumin retention by surfaces appears to be closely related to their surface free energy (in air). Low energy surfaces (in air), whether untreated or glow discharge-treated, retain a larger fraction of the albumin adsorbed than higher energy surfaces. The lowest energy surfaces should have the highest interfacial energies in water, with correspondingly high driving forces for adsorption of

proteins. This can lead to the formation of multiple binding sites upon adsorption, permitting strong hydrophobic interactions, which lead to the observed strong binding.

- L15 ANSWER 24 OF 32 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. DUPLICATE
- AN 1991:27062 BIOSIS
- DN BA91:16413
- TI APPLICATION OF A THERMALLY-REVERSIBLE POLYMER-ANTIBODY CONJUGATE IN A NOVEL MEMBRANE-BASED IMMUNOASSAY.
- AU MONJI N; COLE C-A; TAM M; GOLDSTEIN L; NOWINSKI R C; HOFFMAN A S
- CS GENETIC SYSTEMS CORP., 3005 FIRST AVE., SEATTLE, WASHINGTON 98121.
- SO BIOCHEM BIOPHYS RES COMMUN, (1990) 172 (2), 652-660. CODEN: BBRCA9. ISSN: 0006-291X.
- FS BA; OLD
- LA English
- We have developed a novel method to immobilize antibodies onto a cellulose acetate membrane using a conjugate of an N-isopropylacylamid polymer covalently bound to the antibody. When compared with the unconjugated antibody, over 30-fold increase in retention of the antibody on the membrane was observed when it was conjugated to poly (N-isopropylacrylamide). Studies of the polymer-membrane interaction suggest a combination of hydrophobic and ionic forces, especially the former, is responsible for the high retention. We applied this novel immobilization technology in the development of a membrane-based immunoassay.
- L15 ANSWER 25 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 1989:4185 HCAPLUS
- DN 110:4185
- TI Polymerization induced separation assay using recognition pairs
- IN Thomas, Elaine K.; Schwartz, Dennis E.; Priest, John H.; Nowinski, Robert C.; Hoffman, Allan S.
- PA Genetic Systems Corp., USA
- so U.S., 43 pp.
  - CODEN: USXXAM
- DT Patent
- LA English
- FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
				4 0 0 4 0 6 0 0

- PI US 4749647 A 19880607 US 1984-623838 19840622
- Methods and compds. are disclosed for detg. the presence, amt. of, or AΒ assocn. between substances of interest in samples suspected of contg. same. The methods are based on the polymn.-induced sepn. of specifically bound, reporter-labeled recognition reactants from free, reporter-labeled recognition reactants. The methods described are applicable to any substance for which suitable recognition reactants exist or can be made (e.g. antigen/antibody, hormone/receptor, drug/receptor, nucleic acids, chelating agent/ion, etc.) and are not limited by considerations such as chem. compn. or mol. size. Acrylic acid monomer was conjugated to mouse monoclonal antibody to human IgM .kappa. chains via a spacer arm of p-aminobenzoic acid and a 2nd monoclonal antibody to human IgM .mu. chains was labeled with FITC. A simultaneous sandwich immunoassay for human IqM involved incubating the antibodies with sample, copolymg. with 2-hydroxyethyl methacrylate [initiated with TEMED and (NH4)2S2O8], and analyzing by flow cytometry. The fluorescence intensity of copolymer particles formed in the presence of IgM increased 300-fold over control. The increase in intensity was a linear function of the amt. of IgM present in the sample.

- ANSWER 26 OF 32 HCAPLUS COPYRIGHT 2002 ACS L15
- 1970:438391 HCAPLUS AN
- 73:38391 DN
- Crosslinked poly(hydroxyethyl methacrylate) membranes for TIdesalination by reverse osmosis
- Jadwin, T. A. Hoffman, Allan Sachs; Vieth, W. R. ΑU
- Dep. of Chem. Eng., Massachusetts Inst. of Technol., Cambridge, Mass., USA ÇŞ
- J. Appl. Polymassci. (1970), 14(5), 1339-59 SO CODEN: JAPNAB
- DTJournal
- English LA
- Crosslinked poly(hydroxyethyl methacrylate) membranes for reverse osmosis AB desalination were prepd. by adding trimethylolpropane trimethacrylate (I) or ethylene glycol dimethacrylate to thin film homopolymers. Reverse osmosis, osmosis, and sorption tests were performed. The reverse osmosis H2O flux (at 1500 psi applied pressure and 4% NaCl at pH = 5) of the membranes decréased from 0.6 to 0.055 gal/mils/ft2 day, and the salt rejection increased from 78 to 94% max. as the I concn. increased (0-11 mole %). The #20 content decreased from 42 to 15% over the same I range, but the preferential sorption of H2O to salt did not vary. Rises in reverse-osmosis semipermeability were caused by H2O-NaCl diffusivity ratio changes. A mechanism of permselectivity, in terms of parallel diffusive fluxes across the membrane of primary H-bonded H2O and secondary H2O plus salt ions, is discussed.
- ANSWER 27 OF 32 HCAPLUS COPYRIGHT 2002 ACS L15
- 1970:122509 HCAPLUS AN
- DN 72:122509
- Polyacrylic desalination membranes. II. Reverse osmosis ΤI performance
- ΑU Hoffman, Allan Sachs; Modell, Michael; Pan, Peter
- Dep. of Chem. Eng., Massachusetts Inst. of Technol., Cambridge, Mass., USA CS
- J. Appl. Polym. Sci. (1970), 14(2), 285-301 SO CODEN: JAPNAB
- DTJournal
- LA English
- A new class of polyacrylic membranes was tested under reverse osmosis conditions on dil (1-4%) salt solns. Fluxes up to 0.2 gal-mil/ft2-day at AB >98% rejection have been achieved. The effect of membrane compn. on product flux and salt rejection is discus sed. Increased fluxes at even higher rejection should be possible by proper sel ection of the type and concn. of hydrophilic, hydrophobic, and crosslinkin g monomers. Membranes should have as high as possible a concn. of hydrophilic groups, distributed randomly through a lightly crosslinked, rubbery polymer matrix.
- ANSWER 28 OF 32 HCAPLUS COPYRIGHT 2002 ACS 1969:528543 HCAPLUS L15
- AN
- DN 71:128543
- Polyacrylic desalination membranes. I. Synthesis and ΤI characterization
- Hoffman, Allan Sachs; Modell, Michael; Pan, Peter ΑU
- Massachusetts Inst. of Technol., Cambridge, Mass., USA CS
- J. Appl. Polym, Sci. (1969), 13, 2223-34 SO CODEN: JAPNAB
- DT Journal
- LA English
- AB Polymn. of a mixt. of hydrophilic monomers (N-methylolacrylamide and CH2:CHCO2H), a hydrophobic monomer (CH2:-CHCO2Et), and a hydrophobic

crosslinking monomer (trimethylolpropane trimethacrylate), followed by heat treatment yielded new homogeneous desalination membranes .apprx.6 mils thick. They were characterized by measuring H2O contents and salt distribution coeffs. using an immersion technique. The fractional H2O content in the membrane was 0.16-0.44 with respect to the molal salt distribution coeffs. .apprx.0.22-0.43. A model of intrapolymer H2O is presented: primary H2O is H-bonded with a hydrophilic polymer group while secondary H2O is imbibed with NaCl from the external soln. into hydrophilic regions or defects within the polymer matrix. All compns. contained .apprx.2-3 moles primary H2O/mole hydrophilic monomer. By varying the membrane compn. the sorption characteristics are controlled and can lead to control of flux and permselectivity.

- L15 ANSWER 29 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 1969:461922 HCAPLUS
- DN 71:61922
- TI Structure-property relations for liquid transport in modified polypropylene membranes
- AU Michaels, Alan S.; Vieth, Wolf R.; Hoffman, Allan Sachs; Alcalay, Haim A
- CS Massachusetts Inst. of Technol., Cambridge, Mass., USA
- SO J. Appl. Polym Sci. (1969), 13, 577-98 CODEN: JAPNAB
- DT Journal
- LA English
- AB The permeation and permselective properties of polypropylene (I) films towards org. ligs. and vapors were examd. using films subjected to solvent and thermal treatments. The effect of the treatments on film morphology and transport properties was also detd. and structure-property relations for membrane permeation were developed. I film (Profax 6520F) with 95% isotacticity and a mol. wt. of 3 .times. 105 was extruded onto a casting roll at 100.degree. as a polymer melt to give unoriented hot-cast 5-mil thick films used in the expts. Unoriented quenched films prepd. on a casting roll at 20 degree. were also used. The solvents used were isooctane, methylcyclohexane, PhMe, p-xylene, and o-xylene with soly. parameter differences with respect to I of 1, 0.3, 0.8, 0.6, and 0.9 resp. The I film was solvent modified by immersion in solvent baths at 60-100.degree. for 24 hrs. and samples were dried in vacuo at 40.degree.. Lig. permeation fluxes were detd. using a permeation cell in a thermostatted air bath to prevent concn. of the permeant. A normalized flux rate was calcd. The kinetics of vapor sorption were detd. using a quartz-spring balance at const. temp. by admitting a vapor to the evacuated sorption column at satd. vapor pressure and detg. the spring displacement. Film sample d. was detd. using iso-PrOH-H2O d. gradient columns and used to calc. amorphous vol. fraction. Optical and electron microscope examn. were carried out and melt behavior was observed on a differential scanning calorimeter. The measurements were used to det. permeation, sorption, diffusion, and selectivity in treated and untreated films, the effects of permeation temp. and solvent treatments, and the time dependent vapor transport in I. I films were selective towards PhMe relative to isooctane and p-xylene relative to o-xylene. Liq. flux rates depended on the soly. of the permeant in the films, and the abs. difference in the soly. parameters of the polymer-liq. pair provided a good basis for correlation of this effect. In liqs. with similar soly. parameters, fluxes depended on the apparent mol. cross-section of the permeants. Annealed films showed enhanced permeability but reduced selectivity. These effects resulted from the influence of the solvent type on the polymer morphology and the formation of opened spherulitic structures as a result of recrystn. in the presence of the solvent during annealing. Enhanced flux rates resulted from changes in the spherulite

textures and diminished intercryst. tie chain constrainment within the spherulitic substructure.

- L15 ANSWER 30 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 1969:58483 HCAPLUS
- DN 70:58483
- TI Development of ultrathin skin membranes-hema polymers
- AU Hoffman, Allan S.; Modell, Michael; Hunter, Jack A.; Gillam, W. Sherman; Podall, Harold E.
- CS Massachusetts Inst. of Technol., Cambridge, Mass., USA
- SO U. S. Office Saline Water, Res. Develop. Progr. Rep. (1968), No. 374, 30 pp. Avail.: GPO, 55 cents CODEN: XISWAP
- DT Report
- LA English
- AB A membrane is prepd. by treating a mixt. of acrylic acid 22.7, N-methylolacrylamide 12.3, Et acrylate 40.9, trimethylolpropane trimethacrylate (I) 13.6, and H2O 10.5 vols. with 1% Bz2O2 and a small amt. (2 drops/5 ml. of soln.) of PhNMe2, shaking the compn. for a few sec., pouring it onto Teflon, covering it with glass for 5 min., removing the glass contg. the adherent film, heating the film at 80.degree. for 20 min., and immersing the glass in H2O to release the film, which was 6-8 mils thick and had good mech. properties. This membrane gave slightly better water desalination than did a dense cellulose acetate (39.8% acetylated) membrane. Other membranes prepd. as described above but with smaller amts. of Et acrylate, with no I, or with acrylamide in place of Et acrylate gave less satisfactory desalination. The theory that predicted that the membrane prepd. as described above would be useful in water desalination is discussed.
- L15 ANSWER 31 OF 32 HCAPLUS COPYRIGHT 2002 ACS
- AN 1970:512832 HCAPLUS
- DN 73:112832
- TI Expanded glassy polymers and polyelectrolyte complexes as reverse osmosis and ion-selective membranes
- AU Baddour, Raymond F.; Vieth, Wolf R.; Douglas, Allan S.; Hoffman, Allan S.
- CS Office of Saline Water, Washington, D. C., USA

- SO U.S. Clearinghouse Fed. Sci. Tech. Inform., PB Rep. (1967), No. 191232, 82 pp. Avail.: CFSTI From: U.S. Govt. Res. Develop. Rep. 1970, 70(13), 80 CODEN: XCCRAO
- DT Report
- LA English
- The results of the investigation showed that by varying the membrane AB prepn. conditions it is possible to control the permeability of the film to dissolved salts. For a given material, however, redn. of the salt permeability usually coincides with redn. of the water flux through the membrane. Many of the materials studied in this investigation show very high salt rejection. For example, cellulose nitrate and hydroxyethyl methacrylate (HEMA) both show salt rejections in excess of 90% at low flow rates, while polyurethanes are just below that value for salt rejection. For each of these 3 materials new techniques were developed to control the water flux and salt rejections, and the technology is now available to control the properties of a variety of hydrophilic polymers. The most promising materials for use in reverse osmosis developed on this program appear to be polyurethanes and HEMA. Both of these, however, still must be prepd. as ultrathin films. Other significant results of this investigation have been the development of new characterization techniques to more fully describe polymer properties and to predict performance.

- ANSWER 32 OF 32 HCAPLUS COPYRIGHT 2002 ACS 1968:459874 HCAPLUS L15
- ΑN
- DN 69:59874
- ΤI Expanded glassy polymers and polyelectrolyte complexes as reverse-osmosis and ion-selective membranes
- Baddour, Raymond F.; Vieth, Wolf R.; Douglas, Allan S.; Hoffman, ΑU Allan S.; Hunter, J. A.; Gillam, W. Sherman; Podall, H. E.
- CS Massachusetts Inst. Technol., Cambridge, Mass., USA
- U. S. Dep. Interior, Office Saline Water Res. Develop. Progr. Rep. (1967), SO No. 274, 79 pp. Avail.: GPO, 45 cents CODEN: XISWAP
- DT Report
- LA English
- AB Various classes of semipermeable membranes were tested in programs for developing new materials with higher water permeabilities and longer useful lifetimes than cellulose acetate, while retaining the salt rejection capacity of the latter polymer. Polyurethanes prepd. from tolylene diisocyanates and polyethylene glycol and cross-linked with trimethylolpropane were studied. The water flux through membranes from this material depended only slightly on chem. crosslink d., but showed a strong dependence on the mol. wt. of the I monomer component. The polyelectrolyte complexes from poly(Na styrenesulfonate) and poly(vinylbenzýltrimethylammonium chloride) showed water flux which varied linearly with the effective pressure. The salt flux through the membrane was linear with respect to osmotic pressure differential, and the normalized salt flux was independent of pressure and upstream salt concn. The transport of both salt and water was about to be primarily diffusive in nature, with the contribution of pore flow or hydrodynamic flow through pin holes being small. Poly(.beta.-hydroxyethyl methacrylate) crosslinked with trimethylolpropane trimethacrylate (II) or ethylene glycol, and methacrylic acid-II-.beta.-hydroxyethyl methacrylate copolymers were also tested. The best results obtained were 94% salt rejection at normalized water-flux 0.1 gallon-mil/ft.2-day for II crosslinked with .apprx.9 mole % III. Salt distribution coeffs. for these membranes were comparable to those in cellulose acetate. A selective annealing method for forming skins on expanded cellulose nitrate films was also developed.

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FILE COVERS 1907 - 7 Mar 2002 VOL 136 ISS 10 FILE LAST UPDATED: 5 Mar 2002 (20020305/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

The P indicator for Preparations was not generated for all of the CAS Registry Numbers that were added to the CAS files between 12/27/01 and 1/23/02. As of 1/23/02, the situation has been resolved. Searches and/or SDIs in the H/Z/CA/CAplus files incorporating CAS Registry Numbers with the P indicator executed between 12/27/01 and 1/23/02 may be incomplete. See the NEWS message on this topic for more information. 'OBI' IS DEFAULT SEARCH FIELD FOR 'HCAPLUS' FILE

=> d his

(FILE 'HOME' ENTERED AT 09:54:09 ON 07 MAR 2002)

```
FILE 'HCAPLUS ENTERED AT 09:54:14 ON 07 MAR 2002
L1
            4072 S MEMBRANE? (L) (DISRUPT? OR ALTER?)
L2
          988484 S POLYMER##
L3
           80442 S CONJUGAT?
L4
               4 S L1 AND L2 AND L3
L5
              64 S L1 AND L2
           33942 S HYDROPHO?
L6
L7
           22771 S HYDROPHIL?
          1 S LS AND L6 AND L7
85827 S CELL MEMBRANE/CW
^{18}
L9
              39 S L9 AND L2 AND L3
L10
               5 S L 10 AND (L6 OR L7)
L11
               9 S LS AND (L6 OR L7)
L12
L13
              15 S 1 OR L11 OR L12
```

FILE 'HCAPLUS" ENTERED AT 09:58:46 ON 07 MAR 2002

=> d .ca 1-15

L13 ANSWER 1 OF 15 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:638755 HCAPLUS

TITLE:

Concentration and removal of endocrine

disruptors through hydrophobic

polymeric membrane by pervaporation AUTHOR(S):

Asano, Takao; Yoon, Boo Ok; Hara, Mariko; Higuchi,

CORPORATE SOURCE:

Department of Applied Chemistry, Seikei University,

Tokyo, 180-8633, Japan

SOURCE:

Abstracts of Papers, 222nd ACS National Meeting,

Chicago, IL, United States, August 26-30, 2001 (2001), ENVR-176. American Chemical Society: Washington, D.

CODEN: 69BUZP

DOCUMENT TYPE:

Conference; Meeting Abstract

LANGUAGE:

English

Endocrine disrupting chems., such as dioxin and polychlorinated biphenyl AB (PCB), are affecting the development and reprodn. of humans and animals, and are, therefore, of major concern to the environment. In this work, we examd. the feasibility of removing endocrine disrupting chems. from extremely dil. aq. solns. through hydrophobic polydimethylsiloxane (PDMS) membranes by pervaporation. 1,2-Dibromo-3-chloropropane (DBCP), diethylphthalate, dioxin and biphenyl were selected as model endocrine disrupting chems. The endocrine disrupting chems. could be sepd. very efficiently from dil. aq. solns. through PDMS membranes by pervaporation when the vacuum line between pervaporation cell and a cold trap on the permeate side was heated to 150 -C. The sepn. factors of the endocrine disrupting chems. could not be correlated well with their mol. size. hydrophobic endocrine disrupting chems. showed higher sepn. factors than those of hydrophilic endocrine disrupting chems. using hydrophobic PDMS membranes.

L13 ANSWER 2 OF 15 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER:

2001:528152 HCAPLUS

DOCUMENT NUMBER:

135:261910

TITLE:

Concentration and removal of endocrine

disruptors through hydrophobic

polymeric membranes by pervaporation

AUTHOR(S):

Asano, Takao; Yoon, Boo Ok; Hara, Mariko; Higuchi,

Akon

CORPORATE SOURCE:

Department of Applied Chemistry, Seikei University,

Musashino, Tokyo, 180-8633, Japan

SOURCE:

Preprints of Extended Abstracts presented at the ACS National Meeting, American Chemical Society, Division of Environmental Chemistry (2001), 41(2), 227-232

CODEN: PEACF2; ISSN: 1524-6434

PUBLISHER:

American Chemical Society, Division of Environmental

Chemistry

DOCUMENT TYPE:

Journal; (computer optical disk)

LANGUAGE:

English

AΒ Endocrine disrupting chems. affect human and animal development and reprodn. and are thus of major environmental concern. The feasibility of removing endocrine disrupting chems. from extremely dil. aq. solns. through hydrophobic polydimethylsiloxane (PDMS) membranes via pervaporation was examd. 61-5 (Water)

CC

water purify pervaporation removal endocrine disrupting compd; ST poldimethylsiloxane membrane pervaporation removal endocrine disrupting compd

IT Organic compounds, processes

RL: ADV (Adverse effect, including toxicity); PEP (Physical, engineering

```
or chemical process); POL (Pollutant); REM (Removal or disposal); BIOL
     (Biological study); OCCU (Occurrence); PROC (Process)
        (endocrine disrupting; pervaporation concn. and removal of
        endocrine disrupting compds. through hydrophobic
        polydimethylsiloxane membranes)
ΙT
     Water purification
        (membrane sepn.; pervaporation concn. and removal of
        endocrine disrupting compds. through hydrophobic
        polydimethylsiloxane membranes)
TT
     Membranes, nonbiological
     Water purification
        (pervaporation; pervaporation concn. and removal of endocrine
        disrupting compds. through hydrophobic
        polydimethylsiloxane membranes)
IΤ
     84-66-2, Diethylphthalate
                                  92-52-4, Biphenyl, processes
                                                                   96-12-8,
     1,2-Dibromo-3-chloropropane
                                    104-51-8, N-Butylbenzene
                                                                 262-12-4,
     Dibenzo-p-dioxin 3766-81-2, 2-sec-Butylphenyl methylcarbamate
     22781-23-3, 2,2-Dimethyl-1,3-benzodioxol-4-yl methylcarbamate
     RL: ADV (Adverse effect, including toxicity); PEP (Physical, engineering
     or chemical process); POL (Pollutant); REM (Removal or disposal); BIOL
     (Biological study); OCCU (Occurrence); PROC (Process)
        (pervaporation concn. and removal of endocrine disrupting
        compds. through hydrophobic polydimethylsiloxane
        membranes)
ΙT
     9016-00-6, Polydimethylsiloxane
     RL: DEV (Device component use); TEM (Technical or engineered material
     use); USES (Uses)
        (pervaporation membrane; pervaporation concn. and removal of
        endocrine disrupting compds. through hydrophobic
        polydimethylsiloxane membranes)
REFERENCE COUNT:
                          5
                                THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L13 ANSWER 3 OF 15 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                          2001:525957 HCAPLUS
DOCUMENT NUMBER:
                          135:127195
TITLE:
                          Enhanced transport of therapeutic and diagnostic
                          agents using membrane disruptive
                          acid-sensitive polymers
INVENTOR (S):
                          Hoffman, Allan S.; Stayton, Patrick S.; Murthy, Niren
                          University of Washington, USA
PATENT ASSIGNEE(S):
SOURCE:
                          PCT Int. Appl., 50 pp.
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                       KIND
                             DATE
                                             APPLICATION NO.
                                                               DATE
     WO 2001051092
                       A2
                             20010719
                                             WO 2001-US356
                                                               20010105
     WO 2001051092
                       A3
                             20011206
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
                    CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             CR, CU;
             HU, ID,
             LU, ĽV;
                     MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
                     SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
             SD, SE;
             ZA, ZW;
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
```

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DE, DK; ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                        US 2000-174893
PRIORITY APPLN. INFO.:
                                                         P 20000107
     Compns. and methods for transport or release of therapeutic and diagnostic
     agents, metabolites or other analytes from cells, compartments within
     cells, or through cell layers or barriers are described. The compns.
     include a membrane barrier transport enhancing agent and are usually
     administered in combination with an enhancer and/or exposure to stimuli to
     effect disruption or altered permeability, transport or release. In a
     preferred embodiment, the compns. include compds. which disrupt endosomal
     membranes in response to the low pH in the endosomes but which are
     relatively inactive toward cell membranes (at physiol. pH, but can become
     active toward cell membranes if the environment is acidified below pH
     6.8), coupled directly or indirectly to a therapeutic or diagnostic agent.
     Other disruptive agents can also be used, responsive to stimuli and/or
     enhancers other than pH, such as light, elec. stimuli, electromagnetic
     stimuli, ultrasound, temp., or combinations thereof. The compds. can be
     coupled by ionic, covalent or H bonds to an agent to be delivered or to a
     ligand which forms a complex with the agent to be delivered. Agents to be
     delivered can be therapeutic and/or diagnostic agents. Treatments which
     enhance delivery such as ultrasound, iontophoresis, and/or electrophoresis
     can also be used with the disrupting agents. For example, a terpolymer of
     dimethylaminoethyl methacrylate, Bu methacrylate, and styrene benzaldehyde
     was prepd. for the membrane-disruptive backbone which was then PEGylated
     with thiol-terminated monofunctional or heterofunctional PEGs.
     acid-degradable linkage was a p-aminobenzaldehyde acetal.
IC
     ICM A61K047-48
CC
     63-6 (Pharmaceuticals)
ST
     polymer cell membrane disruption diagnostic
     therapeutic transport
TT
     Diagnosis
        (agents; enhanced transport of therapeutic and diagnostic agents using
        membrane disruptive acid-sensitive polymers
IT
     Polymers, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (conjugates; enhanced transport of therapeutic and diagnostic
        agents using membrane disruptive acid-sensitive
        polymers)
ΙT
    Animal cell
     Cytoplasm
     Organelle
        (delivery to; enhanced transport of therapeutic and diagnostic agents
        using membrane disruptive acid-sensitive
       polymers)
ΙT
    Amino group
     Carboxyl group
     Cell membrane:
     Drug targeting
     Drugs
     Endocytosis
    Endosome
    Hydroxyl group
     Sulfhydryl group
        (enhanced fransport of therapeutic and diagnostic agents using
        membrane disruptive acid-sensitive polymers
IT
    Antisense oligonucleotides
    Biopolymers
    Carbohydrates, biological studies
```

```
Gene, animal
     Nucleotides, biological studies
     Oligonucleotides
     Peptides, biological studies
     Polyoxyalkylenės, biological studies
     Polysaccharides, biological studies
     Proteins, general, biological studies
     Ribozymes
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (enhanced transport of therapeutic and diagnostic agents using
        membrane disruptive acid-sensitive polymers
ΙT
     Electric charge
     Electromagnetic wave
     Electrophoresis
     Iontophoresis
     Liaht
     Sound and Ultrasound
     рН
        (enhanced transport of therapeutic and diagnostic agents using
        membrane disruptive acid-sensitive polymers
        and exposure to stimuli)
     Polymer degradation
        (hydrolytic, acid; enhanced transport of therapeutic and diagnostic
        agents using membrane disruptive acid-sensitive
       polymers)
ΙT
     Functional groups
        (hydroxyacid; enhanced transport of therapeutic and diagnostic agents
        using membrane disruptive acid-sensitive
       polymers)
     Drug delivery systems
ΙT
        (local and systemic; enhanced transport of therapeutic and diagnostic
        agents using membrane disruptive acid-sensitive
       polymers)
IT
     Biological transport
        (permeation; enhanced transport of therapeutic and diagnostic agents
        using membrane disruptive acid-sensitive
       polymers)
TΤ
     Vinyl compounds, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (polymers / enhanced transport of therapeutic and diagnostic
        agents using membrane disruptive acid-sensitive
       polymers)
    Organic compounds, biological studies
ΙT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (small; enhanced transport of therapeutic and diagnostic agents using
       membrane disruptive acid-sensitive polymers
ΙT
    Drug delivery systems
        (topical; enhanced transport of therapeutic and diagnostic agents using
       membrane disruptive acid-sensitive polymers
ΙT
     63-42-3DP, Lactose, pyridylthioacetalstyrene-methacrylate polymer
    derivs. with with methoxy-PEG-thiols 554-38-1DP, Hexalysine,
    pyridylthioacetalstyrene-methacrylate polymer derivs. with with
    methoxy-PEG-thiols 2321-07-5DP, Fluorescein, pyridylthioacetalstyrene-
    methacrylate polymer derivs. with with methoxy-PEG-thiols
    134874-49-0DP, fluorescein/hexalysine/lactose derivs. of
    pyridylthioacetalstyrene-methacrylate polymers
                                                      282732-40-5DP,
    reaction products with methoxy-PEG-thiol derivs. of
```

```
fluorescein/hexalysine/lactose
     RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
        (enhanced transport of therapeutic and diagnostic agents using
        membrane disruptive acid-sensitive polymers
L13 ANSWER 4 OF 15 HCAPLUS COPYRIGHT 2002 ACS
                         2001:489417 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         135:73701
                         Method for the isolation of hydrophobic
TITLE:
                         proteins using a phase partition system with an
                         affinity polymer
                         Tjerneld, Folke; Sivars, Ulf
INVENTOR(S):
PATENT ASSIGNEE(S):
                         Amersham Pharmacia Biotech AB, Swed.
SOURCE:
                         PCT Int. Appl., 30 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                 ----- Patent ·
                 English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
                                           -----
     -----
                      ____
                            -----
    WO. 2001047947
                      A2
                            20010705
                                           WO 2000-EP13025 20001220
     WO 2001047947
                      A3 20011206
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID; IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                        SE 1999-4803
                                                        A 19991227
    A method for sepg. one or more hydrophobic proteins, for instance membrane
     proteins such as integral membrane proteins, from a mixt. of proteins is
     described. The method is characterized in that said mixt. is partitioned
     in a phase system comprising a micelle-enriched aq. phase (micelle phase)
     and a polymer enriched aq. phase (polymer phase). At least part of the
     polymer of the polymer phase carries an affinity ligand that is capable of
     binding to an affinity structure on at least one of said one or more
     hydrophobic proteins. Escherichia coli membranes, contg.
     genetically-modified cytochrome bo3 ubiquinol oxidase having a histidine
     tag, were solupilized in a pentaethyleneglycol mono-n-dodecyl
     ether/dextran 7500 two-phase system. The membrane protein phase was
     removed and washed 3 times with a pure polymer phase before treatment with
     affinity polymer phase contg. sodium perchlorate and allyldextran
    T150-IDA-Cu(II) metal chelate. ICM C07K001-00
IC
     9-9 (Biochemical Methods)
CC
     Section cross reference(s): 6, 7
ST
    hydrophobic protein sepn phase partition affinity
    polymer; cytochrome bo3 ubiquinol oxidase recombinant membrane
    purifn; allyldextran IDA copper chelate affinity partition membrane
    protein
IT
     Partition
        (affinity; method for isolation of hydrophobic proteins using
        phase partition systems with affinity polymers)
```

```
IT
     Chelates
     RL: BPR (Biological process); NUU (Other use, unclassified); PEP
     (Physical, engineering or chemical process); BIOL (Biological study); PROC
     (Process); USES (Uses)
        (as affinity ligands; method for isolation of hydrophobic
        proteins using phase partition systems with affinity polymers
IT
     Ligands
     RL: BPR (Biological process); NUU (Other use, unclassified); PEP
     (Physical, engineering or chemical process); BIOL (Biological study); PROC
     (Process); USES (Uses)
        (conjugates with polymers, binding to
        hydrophobic proteins; method for isolation of
        hydrophobic proteins using phase partition systems with
        affinity polymers)
IT
     Polymers, biological studies
     RL: BPR (Biological process); NUU (Other use, unclassified); PEP
     (Physical, engineering or chemical process); BIOL (Biological study); PROC
     (Process); USES (Uses)
        (conjugates, with affinity ligands binding to
        hydrophobic proteins; method for isolation of
        hydrophobic proteins using phase partition systems with
        affinity polymers)
IT
     Proteins, specific or class
     RL: BPR (Biological process); PEP (Physical, engineering or chemical
     process); PUR (Purification or recovery); BIOL (Biological study); PREP
     (Preparation); PROC (Process)
        (hydrophobic; method for isolation of hydrophobic
        proteins using phase partition systems with affinity polymers
ΙT
     Detergents
        (ionic; method for isolation of hydrophobic proteins using
        phase partition systems with affinity polymers)
ΙT
     Proteins, specific or class
     RL: BPR (Biological process); PEP (Physical, engineering or chemical
     process); PUR (Purification or recovery); BIOL (Biological study); PREP
     (Preparation); PROC (Process)
(membrane, integral; method for isolation of hydrophobic
        proteins using phase partition systems with affinity polymers
IT
     Proteins, specific or class
     RL: BPR (Biological process); PEP (Physical, engineering or chemical
    process); PUR (Purification or recovery); BIOL (Biological study); PREP
     (Preparation); PROC (Process)
        (membrane imethod for isolation of hydrophobic proteins using
        phase partition systems with affinity polymers)
    Affinity
IT
     Buffers
      Cell membrané
     Ionic strength
     Liposomes
    Micelles
     Partition
     Phase
     Sample preparation
     Temperature
    рΗ
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
IT
     Proteins, general, analysis
```

```
RL: AMX (Analytical matrix); ANST (Analytical study)
        (method for isolation of hydrophobic proteins using phase
        partition (systems with affinity polymers)
     Polymers, uses
ΙT
     Salts, uses
     RL: NUU (Other use, unclassified); PEP (Physical, engineering or chemical
     process); PROC (Process); USES (Uses)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
IT
     Polyoxyalkylenes, reactions
     RL: NUU (Other use, unclassified); PEP (Physical, engineering or chemical
     process); RCT (Reactant); PROC (Process); USES (Uses)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
     Polyoxyalkylenes, biological studies
IT
     RL: BPR (Biological process); NUU (Other use, unclassified); PEP
     (Physical, engineering or chemical process); SPN (Synthetic preparation);
     BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)
        (reaction with copper(II) iminodiacetic acid; method for isolation of
        hydrophobic proteins using phase partition systems with
        affinity polymers)
     Proteins, general, biological studies
ΙT
     RL: BPR (Biological process); PEP (Physical, engineering or chemical
     process); PUR (Purification or recovery); BIOL (Biological study); PREP
     (Preparation); PROC (Process)
        (sepn.; method for isolation of hydrophobic proteins using
        phase partition systems with affinity polymers)
ΙT
     69671-26-7DP genetically-modified with C-terminal histidine tag
     RL: BPN (Biosynthetic preparation); PEP (Physical, engineering or chemical
     process); PUR Rurification or recovery); BIOL (Biological study); PREP
     (Preparation) PROC (Process)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
     9004-54-0D, Dextran T150, allyl derivs., reaction with
IT
                                     14219-31-9D, recation with dextran derivs.,
     copper(II) iminodiacetic acid
     and PEG
     RL: BPR (Biological process); NUU (Other use, unclassified); PEP
     (Physical, engineering or chemical process); BIOL (Biological study); PROC
     (Process); USES (Uses)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
IT
     25322-68-3DP, PEG, reaction with copper(II)iminodiacetic acid
     RL: BPR (Biological process); NUU (Other use, unclassified); PEP
     (Physical, engineering or chemical process); SPN (Synthetic preparation);
     BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
IT
     77-86-1, TRIS buffer
                              110-85-0, Piperazine, uses
                                                            151-21-3, Sodium
                             288-32-4, Imidazole, uses
                                                            540-72-7, Sodium
     dodecyl sulfate, uses
     isothiocyanate 1119-94-4, Dodecyltrimethylammonium bromide MOPS 3055-956, Pentaethylene glycol mono-n-dodecyl ether
                                                                        1132-61-2.
                                                                       5704-04-1,
     Tricine
                7365÷44-8
                            7365-45-9, HEPES
                                                 7558-79-4, Disodium hydrogen
                  7558-80-7, Sodium dihydrogen phosphate
     phosphate
                                                             7601-89-0, Sodium
     perchlorate 7647-14-5, Sodium chloride, uses
                                                        7647-15-6, Sodium
     bromide, uses 9002-93-1, Triton X-100 41444-50-2, Octylglucoside 61012-50-8
                                                  9004-54-0, Dextran T500, uses
                                                  69227-93-6
     RL: NUU (Other use, unclassified); PEP (Physical, engineering or chemical process); PROC (Process); USES (Uses) (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
```

```
25322-68-3, PÉĠ
IT
     RL: NUU (Other use, unclassified); PEP (Physical, engineering or chemical
     process); RCT (Reactant); PROC (Process); USES (Uses)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
                                    7719-09-7, Thionyl chloride
     142-73-4, Iminodiacetic acid
ΙT
                                                                    7758-98-7,
     Copper (II) sulfate, reactions
     RL: RCT (Reactant)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
     142-73-4DP, Iminodiacetic acid, reaction products with chlorinated
IT
     polyoxyethylené
                       27252-69-3DP, reaction products with iminodiacetic acid
     27252-69-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (method for isolation of hydrophobic proteins using phase
        partition systems with affinity polymers)
L13 ANSWER 5 OF 15 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                        2001:310488 HCAPLUS
DOCUMENT NUMBER: |
                         134:331596
TITLE:
                         Polymer-lipid conjugate for fusion
                         of target membranes
INVENTOR(S):
                         Martin, Francis J.; Zalipsky, Samuel
PATENT ASSIGNEE(S):
                         Sequus Pharmaceuticals, Inc., USA
                         U.S., 38 pp., Cont.-in-part of U.S. 5,891,468.
SOURCE:
                         CODEN: USXXAM
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION
                      KIND
     PATENT NO.
                             DATE
                                            APPLICATION NO.
                                                              DATE
     US 6224903
                       В1
                             20010501
                                            US 1998-208684
                                                              19981210
     US 5891468
                             19990406
                                            US 1997-949046
                                                              19971010
                                         US 1996-28269
PRIORITY APPLN. INFO.:
                                                         . P
                                                             19961011
                                         US 1997-949046
                                                         A2 19971010
     A fusogenic liposome compn. for delivering a liposome-entrapped compd.
AB
     into the cytoplasm of a target cell is described. The liposomes have an
     outer surface coating of chem. releasable hydrophilic polymer chains which
     shield hydrophobic polymers on the liposome outer surface. Release of the
     hydrophilic polymer chains exposes the hydrophobic polymers for
     interaction with outer cell membranes of the target cells to promote
     fusion of the liposome with the target cells. Also disclosed is a polymer-lipid conjugate for use in promoting fusion between target
     membranes. The conjugate is composed of a first segment composed of a
     hydrophilic polymer and a second hydrophobic polymer segment. The second
     segment is joined to the first segment by a bond effective to release the
     first segment in response to an existing or an induced physiol. condition.
     Attached to the second segment is a vesicle-forming lipid member.
IC
     ICM A61K009-127
NCL
     424450000
CC
     63-5 (Pharmaceuticals)
     Section cross-reference(s): 35
ST
     liposome fusogenic polymer conjugate lipid targeting
IT
     Functional groups
        (alkoxycarbonyl groups; polymer-lipid conjugate for
        fusion of target membranes)
IT
     Redox potential
        (biol.; polymer-lipid conjugate for fusion of
```

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target membranes)
IT
     Animal virus
         (fusion peptides of; polymer-lipid conjugate for
         fusion of target membranes)
IT
     Polymers, biological studies
     RL: PEP (Physical, engineering or chemical process); THU (Therapeutic
     use); BIOL (Biological study); PROC (Process); USES (Uses)
         (hydrophilic; polymer-lipid conjugate for
         fusion of target membranes)
     Polymers, biological studies
ΙT
     RL: PEP (Physical, engineering or chemical process); THU (Therapeutic
     use); BIOL (Biological study); PROC (Process); USES (Uses)
         (hydrophobic; polymer-lipid conjugate for
         fusion of target membranes)
IT
     Erythrocyte
         (liposomal fusion with; polymer-lipid conjugate for
         fusion of target membranes)
IT
     Drug-delivery systems ---
         (liposomes; polymer-lipid conjugate for fusion of
        target membranes)
ΙT
     Antitumor agents
       Cell membrane
     Disulfide group
     Drug targeting
     Fusion, biological
     Genetic vectors,
     Infection
     Inflammation
     Membrane, biological
     Molecular weight distribution
     Neoplasm
         (polymer-lipid conjugate for fusion of target
        membranes)
ΙT
     Polycarbonates, biological studies
     Polyoxyalkylenes, biological studies
     Polyoxyphenylenes
     Polysulfones, brological studies
     RL: BPR (Biological process); PEP (Physical, engineering or chemical
     process); THU (Therapeutic use); BIOL (Biological study); PROC (Process);
     USES (Uses)
         (polymer-lipid conjugate for fusion of target
        membranes)
     Enzymes, biological studies
IT
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); BIOL (Biological study)
         (signal sequence cleavage by; polymer-lipid conjugate
        for fusion of target membranes)
     Lipids, biological studies
IT
     RL: PEP (Physical, engineering or chemical process); THU (Therapeutic
     use); BIOL (Biólogical study); PROC (Process); USES (Uses)
         (vesicle-forming; polymer-lipid conjugate for
        fusion of target membranes)
ΙT
     Peptides, biological studies
     RL: BPR (Biological process); PEP (Physical, engineering or chemical process); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
         (viral fusion; polymer-lipid conjugate for fusion
        of target membranes)
     9001-12-1, Collagenase
9025-39-2, Heparinase
IT
                                9004-06-2, Elastase
                                                        9004-08-4, Cathepsin
                               9040-48-6, Gelatinase
                                                        141907-41-7, Matrix
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metalloproteinase
     RL: BAC (Biological activity or effector, except adverse); BIOL
     (Biological study)
         (polymer-lipid conjugate for fusion of target
        membranes)
IT
     285552-08-1, Plqlwa peptide+ 335596-40-2, Ffgavigtialqvatsagitagiala
                335596-41-3, Fagvviglaalgvataagvtaavalv peptide+
     Fagvvlagaalgvataagitagial peptide+
                                           335596-43-5, Figaiiggvalgvataagit
     peptide+
                335596-44-6, Flgfllgvgsaiasgvavskvlhleg peptide+
     Avgigamflgflgaagstmgaasmtl peptide+
                                            335596-46-8,
     Kftivfphnqkgnwknvpsnyhycps peptide+
                                            335596-47-9,
     Kfpiytildklgpwspidihhlscpn peptide+
                                            335596-48-0,
     Lfgaiagfiengwegmidgwygfrhq peptide+
                                            335596-49-1,
     Ffgaiagfleggwegmiagwhgytsh peptide+
                                            335596-51-5,
     Ifqiddliiqllfvaivetgiggyll peptide+
     RL: BPR (Biological process); PEP (Physical, engineering or chemical
     process); PRP (Properties); THU (Therapeutic use); BIOL (Biological
     study); PROC (Process); USES (Uses)
         (polymer-lipid conjugate for fusion of target
        membranes)
IT
     9002-88-4, Polyethylene
                                9003-07-0, Polypropylene
                                                            9003-53-6,
     Polystyrene 25190-06-1, Polytetramethylene ether
                                                            25322-69-4,
     Polypropylene oxide
     RL: BPR (Biological process); PEP (Physical, engineering or chemical
     process); THU A Therapeutic use); BIOL (Biological study); PROC (Process);
     USES (Uses)
        (polymer-lipid conjugate for fusion of target
        membranes)
ΙT
     59-30-3DP, Folic acid, conjugates
                                          66-72-8DP, Pyridoxal,
     conjugates
     RL: PNU (Preparation, unclassified); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (polymer-lipid conjugate for fusion of target
        membranes)
IT
     9046-10-0
                 80506-64-5
     RL: RCT (Reactant)
        (polymer-lipid conjugate for fusion of target
        membranes)
IT
     179761-24-1P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (polymer-lipid conjugate for fusion of target
        membranes)
IT
     207287-03-4P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (polymer-lipid conjugate for fusion of target
        membranes)
                                THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                         11 .
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L13 ANSWER 6 OF 15 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:284998 HCAPLUS
DOCUMENT NUMBER:
                          133:105858
TITLE:
                          Facilitated transport of organics of biological
                          interest I. A new alternative for the
                          separation of amino acids by fixed-site crown-ether
                         polysiloxane membranes
AUTHOR(S):
                          Barboiu, M.; Guizard, C.; Hovnanian, N.; Palmeri, J.;
                          Reibel, C.; Cot, L.; Luca, C.
CORPORATE SOURCE:
                         Laboratoire des Materiaux et Procedes Membranaires
                         CNRS UMR 5635, Montpellier, F-34296, Fr.
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SOURCE:
                          J. Membr. Sci. (2000), 172(1-2), 91-103
                          CODEN: JMESDO; ISSN: 0376-7388
PUBLISHER:
                          Elsevier Science B.V.
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                          English
     Fixed-site heteropolysiloxane membranes contq. grafted macrocyclic
     receptors can sep. a mixt. of amino acids. These membranes have an
     intermediate configuration between liq. membranes (selective complexation
     by a specific *carrier) and solid membranes (charge interactions). The
     soln.-diffusion model, which was used to analyze exptl. transport results
     for these membranes, provided evidence for a new dual transport
     mechanisms. With an acidic pH of the feed phase, the selectivity of the
     transport (symport of protons) relative to 1-.alpha.-alanine (Ala) was
     high and dependent on the relative hydrophobicity of the amino acids (1-Ph
     aniline (She), leucine (Leu)) (S=7-10), whereas no selectivity was
     obtained when the pH of the feed phase was higher. The proton-driven
     transport increased the flux and caused the transport rates of amino acids
     to be widely spread out due to different mol. recognition principles
     effecting transport. An active transport mechanism of amino acids is
     possibly present in the solid dense polymeric matrix.
CC
     38-3 (Plastics Fabrication and Uses)
     Section cross reference(s): 34
ΙT
     Complexation 5
       Hydrophobicity
     Molecular recognition
     Permeability
     Transport properties
     Zwitterions
        (alternative for sepn. of amino acids by fixed-site
        crown-ether polysiloxane membranes)
IT
     Membranes, nonbiological
        (liq.; alternative for sepn. of amino acids by fixed-site
        crown-ether polysiloxane membranes)
ΙT
     Amino acids, processes
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
        (neat and in zwitterionic form; alternative for sepn. of
        amino acids by fixed-site crown-ether polysiloxane membranes)
IT
     Polysiloxanes, uses
     RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
        (poly(ether sulfonate)-supported; alternative for sepn. of
        amino acids by fixed-site crown-ether polysiloxane membranes)
ΙT
     Polysulfones, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (polyether-, polymeric membranes supported with; alternative for sepn. of amino acids by fixed-site crown-ether
        polysiloxane membranes)
ΙT
     Polyethers, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (polysulfone polymeric membranes supported with;
        alternative for sepn. of amino acids by fixed-site crown-ether
        polysiloxane membranes)
ΙT
                                    56-41-7, L-.alpha.-Alanine, processes
     56-40-6, Glycine, processes
                                     56-84-8, Aspartic acid, processes
     56-45-1, L-Serine, processes
     56-87-1, L-Lysine, processes
                                     61-90-5, L-Leucine, processes
     L-Phenyl alanine, processes
     RL: PEP (Physiçal, engineering or chemical process); PROC (Process)
        (neat and in zwitterionic form; alternative for sepn. of
        amino acids by fixed-site crown-ether polysiloxane membranes)
IT
     283614-51-7
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RL: PRP (Properties); TEM (Technical or engineered material use); USES
     (Uses)
         (poly(ether)sulfonate)-supported; alternative for sepn. of
         amino acids by fixed-site crown-ether polysiloxane membranes)
REFERENCE COUNT:
                                 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS
                           17
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L13 ANSWER 7 OF 15 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                           2000:255871 HCAPLUS
DOCUMENT NUMBER:
                           133:103745
TITLE:
                           Modulation of immobilized enzyme activity by
                           altering the hydrophobicity of
                           nylon-grafted membranes Part 1. Isothermal
                    ۶.
                           conditions
AUTHOR(S):
                           El-Masry, M. M.; De Maio, A.; Di Martino, S.; Diano,
                           N.; Bencivenga, U.; Rossi, S.; Grano, V.; Canciglia,
                           P.; Portaccio, M.; Gaeta, F. S.; Mita, D. G.
CORPORATE SOURCE:
                          International Institute of Genetics and Biophysics of
                           CNR, Naples, 80125, Italy
                    A Chicken
SOURCE:
                           J. Mol. Catal. B: Enzym. (2000), 9(4-6), 219-230
                           CODEN: JMCEF8; ISSN: 1381-1177
                    13.7
                           Elsevier Science B.V.
PUBLISHER:
DOCUMENT TYPE:
                           Journal
LANGUAGE:
                           English
AB
     The catalytic behavior under isothermal conditions of two different
     membranes loaded with .beta.-galactosidase was investigated. One membrane
     (M1) was constituted by a nylon sheet grafted with methylmethacrylate by
     means of chem grafting. The other, (M2), was prepd. by a double chem. grafting: the first one with styrene (Sty) and the second one with
     methylmethacrylate. Membrane activity was characterized as a function of
     temp., pH and substrate concn. The role of Sty in increasing membrane
     hydrophobicity has been discussed. Membrane M2 was found to be better
     suited for employment in non-isothermal bioreactors.
CC
     16-1 (Fermentation and Bioindustrial Chemistry)
     Section cross-reference(s): 7, 9 Immobilization, biochemical
ΙT
         (enzyme; modulation of immobilized enzyme activity under isothermal
        conditions by altering the hydrophobicity of
        nylon-grafted membranes)
ΙT
     Polyamides, preparation
     RL: NUU (Other use, unclassified); PNU (Preparation, unclassified); PREP
     (Preparation); USES (Uses)
         (graft polymers; modulation of immobilized enzyme activity
        under isothermal conditions by altering the
        hydrophobicity of nylon-grafted membranes)
     Polyamide fibers, uses
RL: NUU (Other use, unclassified); USES (Uses)
IT
        (membrane grafted with methylmethacrylate or styrene and methylmethacrylate; modulation of immobilized enzyme activity under
        isothermal conditions by altering the hydrophobicity
        of nylon-grafted membranes)
IT
     Enzyme kinetics
       Hydrophobicity
     Temperature effects, biological
         (modulation) of immobilized enzyme activity under isothermal conditions
        by altering the hydrophobicity of nylon-grafted
        membranes)
IT
     50-99-7P, Dextrose, preparation
     RL: BPN (Biosynthetic preparation); BIOL (Biological study); PREP
```

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(modulation) of immobilized enzyme activity under isothermal conditions
        by altering the hydrophobicity of nylon-grafted
IT
     9031-11-2, .beta.-Galactosidase
     RL: BPR (Biological process); CAT (Catalyst use); BIOL (Biological study);
     PROC (Process); USES (Uses)
        (modulation of immobilized enzyme activity under isothermal conditions
        by altering the hydrophobicity of nylon-grafted
        membranes)
ΙT
     63-42-3, Lactose
     RL: BPR (Biological process); RCT (Reactant); BIOL (Biological study);
     PROC (Process)
        (modulation of immobilized enzyme activity under isothermal conditions
        by altering the hydrophobicity of nylon-grafted
        membranes)
     124-09-4, Hexamethylenediamine, reactions
                                                 9003-53-6D, PolyStyrene,
ΙT
     grafted copolymer-with nylon membrane and polymethylmethacrylate
     9011-14-7D, PolyMethylmethacrylate, grafted copolymer with nylon
     membrane
     RL: NUU (Other use, unclassified); RCT (Reactant); USES (Uses)
        (modulation of immobilized enzyme activity under isothermal conditions
        by altering the hydrophobicity of nylon-grafted
        membranes)
                         44
                               THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L13 ANSWER 8 OF 15 HCAPLUS COPYRIGHT 2002 ACS
                         1999:451212 HCAPLUS
ACCESSION NUMBER:
                         131:106813
DOCUMENT NUMBER:
                         Enhanced transport using membrane
TITLE:
                         disruptive agents
                         Hoffman, Allan S.; Stayton, Patrick; Press, Oliver;
INVENTOR(S):
                         Tirrell, David; Murthy, Niren; Lackey, Chantal; Crum,
                         Lawrence A.; Mourad, Pierre D.; Porter, Tyrone M.
PATENT ASSIGNEE(S)
                         University of Washington, USA; University of
                         Massachusetts
                         PCT Int. Appl., 54 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION
     PATENT NO.
                      KIND
                            DATE
                                           APPLICATION NO.
                                                            DATE
                                           _____
     WO 9934831
                       A1
                            19990715
                                           WO 1999-US122
                                                            19990105
         W: AU, CA
                     JP.
                    CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
         RW: AT, BE,
             PT, SE
     AU 9920261
                            19990726
                                           AU 1999-20261
                                                            19990105
                       A1
                                           EP 1999-900750
     EP 1044021
                      A1
                            20001018
                                                            19990105
             AT, BE CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, ĘŪį
                            20010712
     US 2001007666
                       A1
                                           US 1999-226044
                                                            19990105
     JP 2002500201 T2
                            20020108
                                           JP 2000-527278
                                                            19990105
PRIORITY APPLN. INEO .:
                                        US 1998-70411
                                                         Ρ
                                                            19980105
                                        WO 1999-US122
                                                         W
                                                            19990105
AB
     Compns. and methods for transport or release of therapeutic and diagnostic
     agents or metabolites or other analytes from cells, compartments within
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cells, or through cell layers or barriers are described. The compns.
include a membrane barrier transport enhancing agent and are usually
administered in combination with an enhancer and/or exposure to stimuli to
effect disruption or altered permeability, transport or release. In a
preferred embodiment, the compns. include compds. which disrupt endosomal
membranes in response to the low pH in the endosomes but which are
relatively inactive toward cell membranes, coupled directly or indirectly
to a therapeutic or diagnostic agent. Other disruptive agents can also be
used, responsive to stimuli and/or enhancers other than pH, such as light,
elec. stimuli, electromagnetic stimuli, ultrasound, temp., or combinations
          The compds. can be coupled by ionic, covalent or H bonds to an
agent to be delivered or to a ligand which forms a complex with the agent
to be delivered. Agents to be delivered can be therapeutic and/or
diagnostic agents. Treatments which enhance delivery such as ultrasound,
iontophoresis, and/or electrophoresis can also be used with the disrupting
agents. The ability of the GALA peptide to lyse erythrocytes was compared
with that of an GALA/poly(acrylic acid) conjugate at pH 5.0. The
conjugate gave 10%—lysis at 100 .mu.g.
ICM A61K047-32
ICS A61K047-42; A61K047-48; A61K041-00
63-6 (Pharmaceuticals)
Section cross-reference(s): 37
drug transport membrane disruptive agent prepn;
polymer protein conjugate drug transport prepn
Immunoglobulins
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
   (G, conjugates, with polymers; enhanced drug
   transport using membrane disruptive agents)
Biological transport
   (drug; enhanced drug transport using membrane
   disruptive agents)
Drug delivery systems
   (emulsions; enhanced drug transport using membrane
   disruptive àgents)
Cell membrane
Cytotoxic agents
Electric field
Electrophoresis
Endosome
Erythrocyte
Gene therapy
Hemolysis
Iontophoresis
Sound and Ultrasound
   (enhanced drug transport using membrane disruptive
   agents)
Polymer blends
RL: POF (Polymer in formulation); THU (Therapeutic use); BIOL (Biological
study); USES ((Uses)
   (enhanced drug transport using membrane disruptive
   agents)
Lipids, biological studies
Nucleic acids in Nucleosides, prological studies
Nucleotides, bidlogical studies
RL: THU (The table utic use); BIOL (Biological study); USES (Uses)
   (enhanced drug transport using membrane disruptive
   agents)
Drug delivery systems
   (liposomes; enhanced drug transport using membrane
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disruptive agents)
IT
     Drug delivery systems
         (microparticles; enhanced drug transport using membrane
        disruptive agents)
     Drug delivery systems
IT
         (nanoparticles; enhanced drug transport using membrane
        disruptive agents)
                                          9003-01-4D, Poly(Acrylic
IT
     79-10-7D, Acrylic acid, polymers
     acid), protein conjugates
     RL: POF (Polymer in formulation); THU (Therapeutic use); BIOL (Biological
     study); USES (Uses)
        (enhanced drug transport using membrane disruptive
        agents)
IT
     107658-43-5DP, Peptide GALA (synthetic pore-forming), polymer
     conjugates
     RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
     (enhanced drug transport using membrane disruptive
        agents)
     9013-20-1D, Streptavidin, conjugates with polymers
IT
     25119-83-9, Acrylic acid-butyl acrylate copolymer
                                                            62607-09-4,
     Poly(ethacrylic acid)
                             62607-09-4D, Poly(ethacrylic acid), protein
     conjugates
                  75034-36-5, Acrylic acid-propyl acrylate copolymer
     138134-74-4, Poly(.alpha.-propylacrylic acid) 138134-76-6,
     Poly(.alpha.-butylacrylic acid)
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (enhanced drug transport using membrane disruptive
        agents)
                         8
                                 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L13 ANSWER 9 OF 15 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                          1997:740133 HCAPLUS
DOCUMENT NUMBER:
                          128:26910
                          Polypeptide conjugates for transporting
TITLE:
INVENTOR(S):
PATENT ASSIGNEE(S)
                          substances across cell membranes
                          Summerton, James E.; Weller, Dwight D.
                          Antivirals Inc., USA
                         PCT Int. Appl., 72 pp.
                          CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                          English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION
                     . KIND
     PATENT NO.
                             DATE
                                             APPLICATION NO.
                                                               DATE
                 A2
     WO 9740854
                             19971106
                                             WO 1997-US7335
                                                               19970430
         W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK,
             EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT,
             RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN,
             ML, MR, NE, SN, TD, TG
     AU 9729298
                     A1
                             19971119
                                             AU 1997-29298
                                                               19970430
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20010208

19991229

AT, BE, CH, DE, FR, GB, IT, LI, LU, NL

EP 1997-923513

19970430

B2 A2

AU 729643

EP 966303

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JP 2000509394
                             20000725
                       T2
                                             JP 1997-539221
                                                              19970430
PRIORITY APPLN. INFO .:
                                         US 1996-16347
                                                          P 19960501
                                         US 1996-28609
                                                           P 19961023
                                         WO 1997-US7335
                                                           W 19970430
     Polymeric compas. effective for delivering compds. in living organisms are
AB
     described. The compns, include polypeptides which exhibit soly, in both
     hydrophilic and lipophilic environments by undergoing a reversible
     pH-dependent transition from a low-pH, lipophilic form to a high-pH,
     hydrophilic form.
     ICM A61K047-48
IC
     63-5 (Pharmaceuticals)
CC
ST
     transdermal drug delivery peptide conjugate antitumor; antiulcer
     drug delivery peptide conjugate; anticaries drug delivery
     peptide conjugate
ΙT
     Proteins (specific proteins and subclasses)
     RL: PEP (Physical, engineering or chemical process); THU (Therapeutic
     use); BIOL (Biological study); PROC (Process); USES (Uses)
        (conjugates polypeptide conjugates for
        transporting substances across cell membranes)
IT
     Stratum corneum (epidermis)
        (extracellular matrix; polypeptide conjugates for
        transporting substances across cell membranes)
ΙT
     Polymers, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (nucleic acid-binding; polypeptide conjugates for
        transporting substances across cell membranes)
ΙT
     Nucleic acids
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (polymers binding; polypeptide conjugates for
        transporting substances across cell membranes)
IT
     Antibacterial agents
     Antitumor agents
       Cell membrane
     Dentifrices
     Drug delivery systems
     Helicobacter pylori
       Hydrophilicity
     Lipophilicity
     Partition
     Protein sequençes
     Transdermal drug delivery systems
     .alpha.-Helix (protein conformation)
        (polypeptide conjugates for transporting substances across
        cell membranes)
ΙT
     25513-46-6, Polyglutamic acid
     RL: BOC (Biological occurrence); THU (Therapeutic use); BIOL (Biological
     study); OCCU (Occurrence); USES (Uses)
        (polypeptide conjugates for transporting substances across
        cell membranes)
IT
     1397-89-3D, Amphotericin b, polypeptide conjugates
     33069-62-4D, Taxol, polypeptide conjugates Cyclosporin, polypeptide conjugates
                                                    79217-60-0D,
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (polypeptide conjugates for transporting substances across
        cell membranes)
     56-41-7, Alanine, biological studies 56-86-0, studies 61-90-5, Leucine, biological studies
IT
                                              56-86-0, Glutamic acid, biological
                                                        63-68-3, Methionine,
                           107-95-9, .beta.-Alanine
                                                       2835-81-6, 2-Amino butyric
     biological studies
           6600-4054, Norvaline
     RL: BOC (Biological occurrence); BSU (Biological study, unclassified);
```

BIOL (Biological study); OCCU (Occurrence) (polypeptides contg.; polypeptide conjugates for transporting substances across cell membranes)

L13 ANSWER 10 OF 15 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1997:667263 HCAPLUS

DOCUMENT NUMBER: 127:322794

TITLE:

Property-affecting and/or property-exhibiting compositions for therapeutic and diagnostic uses INVENTOR (S): Rabbani, Elazar; Stavrianopoulos, Jannis G.; Donegan, James J.; Liu, Dakai; Kelker, Norman E.; Engelhardt,

Dean L.

PATENT ASSIGNEE (S): ; Enzo Therapeutics, Inc., USA

14 % Can. Pat. Appl., 275 pp. SOURCE:

CODEN: CPXXEB

DOCUMENT TYPE:

Patent English LANGUAGE:

FAMILY ACC. NUM. GOUNT: 1

PATENT INFORMATION:

PA	TENT	NO.		KIND	DATE		API	PLICATI	ON 1	10.	DATE	
			÷									
CA	2190	0304		AA	199706	616	CA	1996-2	21903	304	19961:	114
EP	7793	365	1	A2	199706	518	EP	1996-1	1996	51	199612	212
EP	779:	365		АЗ	199911	124						
	R:	DE,	FR,	GB, IT								
JP	093	13190	1	A2	199712	209	JP	1996-3	36004	13	199612	216
US	200	100683	L4	A1	200107	705	US	1997-9	97863	33	199711	125
US	200	100683	15 👫	A1	200107	705	US	1997-9	7863	34	19971	125
US	200	10068	L6	A1	200107	705	US	1997-9	7863	37	199711	125
US	2003	100776	67	A1	200107	712	US	1997-9	7863	32	19971	125
PRIORIT	Y API	PLN.	INFO.	· •		U	S 199	95-5744	143	Α	199512	215
AB Co	mpns	. use:	Eul 1	or effe	ecting	and/or	exhib	oiting	char	iges	in bid	ol.

Compns. useful! for effecting and/or exhibiting changes in biol. functioning and processing in cells and biol. systems are provided which combine chem. modifications and/or ligand addns. with biol. functions in such a way as hot to interfere substantially with the biol. functions. Such addnl. characteristics include nuclease resistance, targeting specific cells or cell receptors, and augmenting or decreasing interactions between the compns. and target cells. A title compn. may constitute a mucleotide, nucleotide analog, nucleic acid, natural or synthetic polymer, ligand, or conjugate of a ligand with any of the preceding. För example, single-stranded DNA from a plasmid contg. a gene of interest is complexed with an allylamine phosphoramidite-contg. oligonucleotide primer (complementary to a region of the DNA distant from the gene of interest) which as been modified with trilactosyllysyllysine (prepn. given) and the primer is extended with Klenow enzyme to form completely double-stranded DNA. On exposure of target cells to this DNA, the galactose moieties on the DNA bind to receptors on the cells, resulting in transport of the DNA into the cells. In another embodiment, DNA for antisense RNA sequences to regions of the HIV genome were inserted into the U1 small nuclear RNA coding region and the DNA was used to transform U937 cells. The transformed cells were resistant to HIV infection, as shown by inhibition of virus replication and p24 antigen prodn.

IC

ICM C07H021-00 A61K047-48; A61K031-70; A61K038-55

CC 63-6 (Pharmaceuticals)

Section cross-reference(s): 3 polynucleotide conjugation ligand cell targeting; protein ST conjugation ligand cell targeting; HIV gene therapy; biopolymer

```
cell targeting
ΙT
     Bacteria (Eubacteria)
     Eukaryote (Eukaryotae)
     Prokaryote
        (DNA of, conjugates with ligands; property-affecting and/or
        property-exhibiting compns. for therapeutic and diagnostic uses)
IT
     Ligands
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (conjugated, with nucleic acids; property-affecting and/or
        property-exhibiting compns. for therapeutic and diagnostic uses)
IT
        (conjugates with ligands; property-affecting and/or
        property-exhibiting compns. for therapeutic and diagnostic uses)
IT
     Biopolymers
     Fatty acid esters
     RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
       (conjugates with nucleic acids; property-affecting and/or
        property-exhibiting compns. for therapeutic and diagnostic uses)
ΙT
     Polyelectrolytes
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (conjugates with nucleic acids; property-affecting and/or
        property-exhibiting compns. for therapeutic and diagnostic uses)
     Fatty acids, biological studies
ΙT
       Polymers, biological studies
     Proteins (specific proteins and subclasses)
     RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical
     study); BIOL (Brological study); USES (Uses)
        (conjugates with nucleic acids; property-affecting and/or
        property-exhibiting compns. for therapeutic and diagnostic uses)
IT
     Bond
     Molecules
        (hydrophobic: property-affecting and/or property-exhibiting
        compns. for therapeutic and diagnostic uses)
IT
     Cell membrane
     Cytoplasm
        (localization to; property-affecting and/or property-exhibiting compns.
        for therapeutic and diagnostic uses)
ΙT
     Bacteriophage ?
     Viroid
         (nucleic acid of, conjugates with ligands; property-affecting
        and/or property-exhibiting compns. for therapeutic and diagnostic uses)
IT
     Animal virus
        (nucleic acids of, conjugates with ligands;
        property-dffecting and/or property-exhibiting compns. for therapeutic
        and diagnostic uses)
ΙT
     Antibody conjugates
     Monoclonal antibody conjugates
     Polysaccharide conjugates
     RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)
         (with nucleic acids; property-affecting and/or property-exhibiting
        compns. for therapeutic and diagnostic uses)
     9004-10-8DP, Insulin, conjugates with oligo(T)
TT
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (property-affecting and/or property-exhibiting compns. for therapeutic and diagnostic uses)
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HCAPLUS COPYRIGHT 2002 ACS
    ANSWER 11 OF 15
ACCESSION NUMBER:
                          1997:372273 HCAPLUS
DOCUMENT NUMBER:
                          126:347323
                          Buccal delivery of glucagon-like insulinotropic
TITLE:
                          peptides (GLPs)
INVENTOR(S):
                          Heiber, Sonia J.; Ebert, Charles D.; Gutniak, Mark K.
PATENT ASSIGNEE(S):
                          Theratech, Inc., USA
                          PCT Int. Appl., 55 pp.
SOURCE:
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
                          English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                       KIND
                             DATE
                                             APPLICATION NO.
                                                                DATE
                     : ----
                             _____
                                             -----
     _____
                                             WO 1996-US16890 19961022
                        A1
                             19970501
         W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT,
             RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: KE, LS; MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR,
              IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI
                                             US 1995-553807
                              19980616
                                                                19951023
     US 5766620
                        Α
                              19970501
                                             CA 1996-2235369
                                                                19961022
     CA 2235369
                             19970515
                                             AU 1996-74647
     AU 9674647
                        Α1
                                                                19961022
                             20000217
     AU 716038
                        В2
                                             EP 1996-936815
                                                                19961022
                       A1
                             19980826
     EP 859606
             AT, BE CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
                             19981223
                                             CN 1996-198618
                                                                19961022
     CN 1202820
                             19990406
                                             BR 1996-11139
                                                                19961022
     BR 9611139
                        Α
                        Т2
                              19991130
                                             JP 1996-516712
                                                                19961022
     JP 11513982
                        В
                              20010101
                                             TW 1996-85112962 19961022
     TW 416854
                                             ZA 1996-8909
     ZA 9608909
                        A
                             19970528
                                                                19961023
                     ). A
                                             US 1997-964731
     US 5863555
                             19990126
                                                                19971105
                                          US 1995-553807
PRIORITY APPLN. INFO .:
                                                            Α
                                                               19951023
                                          WO 1996-US16890 W 19961022
     Drug delivery systems for administering a GLP to the buccal mucosa for
AB
     transmucosal drug delivery comprise a drug compn. contg. effective amts.
     of the GLP and a permeation enhancer, and means for maintaining the drug
     compn. in a drag-transferring relation with the buccal mucosa. These
     systems can be in free form, such as creams, gels, and ointments, or can
     comprise a device of detd. phys. form, such as tablets, patches, and
     troches. A préferred GLP is GLP-1(7-36) amide. Thus, a gingival bilayer
     tablet was prepd. comprising an active layer and an adhesive layer. The
     adhesive layer was prepd. by mixing polyethylene oxide 70, Carbopol 934P
     20, and compressible xylitol/CM-cellulose filler 10 wt. parts, granulating
     with EtOH, sieving, drying, mixing with stearic acid 0.25 and mint flavor
     0.06 wt.%, and compression. To prep. the active layer, mannitol 49.39,
     hydroxypropylcellulose 34.33, and Na taurocholate 15.00 wt.% were mixed,
     granulated with EtOH, sieved, dried, combined with GLP-1(7-36) amide 0.91, FD&C Yellow No. 6HT 0.06, Mg stearate 0.25, and mint flavor 0.06 wt.%; 50 mg of this mixt, was compressed onto 50 mg adhesive layer.

ICM A61K009-70

ICS A61L015-16
IC
CC
     63-6 (Pharmaceuticals)
IT
     Caseins, biological studies
     Gelatins, biològical studies
```

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Polyethers, biological studies
     Vinyl polymers
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (adhesives contg.; buccal delivery of glucagon-like insulinotropic
     peptides)
Cell membrane
IT
        (disrupting agents for; buccal delivery of glucagon-like
        insulinotropic peptides)
     Polymers, biological studies
IT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (hydrophilic, adhesive contg.; buccal delivery of
        glucagon-like insulinotropic peptides)
ΙT
     79-10-7D, 2-Propenoic acid, esters, polymers
     2-Propenoic acid, polymers 557-75-5D, Ethenol,
                                        9000-69-5, Pectin
     polymers
                 9000-30-0, Guar gum
                                                             9003-39-8,
                        9004-54-0, Dextran, biological studies
                                                                    9004-57-3,
            9004-32-4
     Ethylcellulose
                       9004-62-0, Hydroxyethylcellulose
                                                            9004-64-2,
    Hydroxypropyleellulose 9004-65-3, Hydroxypropylmethylcellulose
     9005-25-8, Starch, biological studies
                                                25322-68-3
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
         (adhesive contg.; buccal delivery of glucagon-like insulinotropic
        peptides)
     107-35-7D, Taurine, bile acid conjugates, salts
                                                          12441-09-7D,
IT
                        25312-65-6D, Cholanic acid, salts
     Sorbitan, esters
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (permeation enhancers; buccal delivery of glucagon-like insulinotropic
        peptides)
L13 ANSWER 12 OF 15 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                          1992:262361 HCAPLUS
                          116:262361
DOCUMENT NUMBER:
                          Modified polyanionic polymers. I: Grafting
TITLE:
                          of hydrophobic group onto poly(maleic
                          acid-alt-3,4-dihydroxyphenylprop-1-ene) to improve the
                          affinity for cell membranes
                          Suda, Yasuo; Yamamoto, Hitomi; Sumi, Masao; Oku,
AUTHOR(S):
                          Naoto; Ito, Fumiaki; Yamashita, Shinji; Nadai,
                          Tanekazu; Ottenbrite, Raphael M.
                          Fac. Sci., Osaka Univ., Toyonaka, 560, Japan J. Bioact. Compat. Polym. (1992), 7(1), 15-24
CORPORATE SOURCE:
SOURCE:
                          CODEN: JBCPEV; ISSN: 0883-9115
DOCUMENT TYPE:
                           Journal
LANGUAGE:
                          English
     To improve the affinity of polyanionic polymers for cell membranes,
AB
     several hydrophobic groups were grafted onto poly(maleic
     acid-alt-3,4-dihydroxyphenylprop-1-ene) [poly(MA-alt-DP)] which has
     cytotoxic activity. The effect of the degree of substitution of the
     grafted group to the maleic anhydride residue was also evaluated. Grafted
     polymers were characterized by their partition coeffs., their affinity to liposomes and the ability to interact with rat small intestinal epithelial
     cells. The cell affinity of the modified polyanionic polymers could be
     augmented and controlled by simple grafting.
     63-5 (Pharmaceuticals)
CC
     Section cross reference(s): 1, 35 maleate dihydroxyphenylpropene polymer amine graft; cell
ST
     affinity polymer
     Cell membrane
IT
         (affinity of amine grafted-diacetoxyphenylpropene-maleic anhydride
        alternating copolymer for)
     Lipophilicity
IT
```

(of amine grafted-diacetoxyphenylpropene-maleic anhydride alternating copolymers, affinity for cell membranes in relation to)

IT Amines, compounds

RL: SPN (Synthetic preparation); PREP (Preparation) (reaction products, with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed, prepn. and affinity for cell membranes of)

IT 62-53-3DP, Aniline, reaction products with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed 107-10-8DP, Propylamine, reaction products with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed 109-73-9DP, Butylamine, reaction products with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed 111-26-2DP, Hexylamine, reaction products with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed 111-86-4DP, Octylamine, reaction products with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed - 2016-57-1DP; Decylamine, reaction products with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed 67247-04-5DP, reaction products with diacetoxyphenylpropene-maleic anhydride alternating copolymer, hydrolyzed 141596-23-8DP, reaction products with amines, hydrolyzed

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and affinity for cell membranes of)

IT 13620-82-1P

> RL: RCT (Readtant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and polymn. of, with maleic anhydride)

L13 ANSWER 13 OF 15 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1991:602736 HCAPLUS

DOCUMENT NUMBER:

TITLE:

115:202736 Membrane affinity purification apparatus and its use in the purification of macromolecules of therapeutic

value

INVENTOR(S): Goffe, Randal A.; Zale, Stephen E.; O'Connor, James

L.; Kessler, Stephen B.; Cohen, Charles M.

PATENT ASSIGNEE (S)

SOURCE:

Sepracor, Inc., USA PCT Int. Appl., 142 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION: 14

PAT	TENT NO.	1000	KIND	DATE		A	PPLI	CATI	ои и	٥.	DATE				
. WO	9005018		A1	19900517	- 7	W	0 19	 89-บ	5484	7	1989	1030			
	W: AU,	BB,	BG, BI	R, DK, FI,	HŲ,	JP,	KR,	LK,	MC,	MG,	MW,	NO,	RO,	SD,	SU
	RW: AT,	BE,	BF, B	J, CF, CG,	CH,	CM,	DE,	FR,	GΑ,	GB,	IT,	LU,	ML,	MR,	
	NL,	SE,	SN, TI	O, TG										-	
CA	2001720	1 : 4	AA	19900430	)	C.	A 19	39-2	0017	20	1989	1027			
ΑU	8945247	1.1	A1	19900528	}	Α	U 19	39-4	5247		1989	1030			
EΡ	483143		. A1	19920506	5	E	P 19	39-9	1270	2 -	1989	1030			
EΡ	483143	The second	, B1	19940601	L										
EΡ	483143	7.0	B2	19970409	)										
	R: AT,	BÉ,	CH, DI	E, FR, GB,	IT,	LI,	LU,	NL,	SE						
AT	106272	3	, E	19940615	5	Α	T 19	39-9	1270	2	1989	1030			
US	5310688	100	A	19940510	)	Ü	S 19	93-3	5549		1993	0323			
US	5683916		Α	19971104	ļ	U	S 19	95-4	6547	9	1995	0605			

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PRIORITY APPLN. INFO .:
                                                 US 1988-265061
                                                                         19881031
                                                 US 1989-428263
                                                                         19891026
                                                 EP 1989-912702
                                                                         19891030
                                                WO 1989-US4847
                                                                         19891030
                                                 US 1990-487668
                                                                         19900302
                                                 US 1993-83859
                                                                         19930628
     An app. is provided which is useful for the sepn. of .gtoreq.1 preselected
AB
      ligate(s) in a fluid. Also provided is an easily scaled-up membrane
      affinity sepn. process which is reliable, highly selective, gives a high
      yield of product, and has a high volumetric throughput. A substantially
      isotropic porous membrane is used, to which is assocd. a preselected
      ligand, which provides an optimum loading capacity and low dead vol. while
     allowing high filtrate flow rates. Methods for isolation of macromols. of therapeutic value, e.g. factor VIII and fibronectin, are described, and
     diagrams of the app. are included. Cloning and expression of a bifunctional binding site protein (one domain binding digoxin and the
     other binding Ig Fc regions) are also described. Thus a polyether sulfone/poly(ethylene oxide) hollow-fiber membrane was sequentially reacted with ethylene glycol diglycidyl ether and hydroxyethyl cellulose,
      activated with 2-fluoro-1-methylpyridinium p-toluenesulfonate, and the
      activated fibers reacted with an antibody to factor VIII. The resulting
     membrane was used to purify a factor VIII conc.; the purifn. factor was
      B01D063-02; B01D063-04; B01D063-08; C07K003-20
IC
     9-3 (Biochemical Methods)
Section crossite ference(s): 63
Blood-coagulation factors
CC
IT
      Interferons
      RL: ANST (Analytical study)
         (antibody to, conjugates with hollow-fiber membrane polymer, for biomol. sepn.)
ΙT
     Liposome
      Plasmid and Episome
      Surfactants
     Agglutinins and Lectins
     Antibodies
     Antigens
     Blood-coagulation factors
     Hormones
     Receptors
     RL: ANST (Analytical study)
          (conjugates with hollow-fiber membrane polymer, for
         biomol. sepn.)
ΙT
      Carboxylic acids, compounds
      RL: ANST (Analytical study)
          (conjugates, with hollow-fiber membrane polymer,
          for biomol. sepn.)
ΙT
     Macromolecular compounds
      RL: ANST (Analytical study)
          (hollow-fiber affinity membrane contg., for biomol. sepn.,
         surface property alteration in relation to)
     Polycarbonates, uses and miscellaneous Polyesters, uses and miscellaneous
IT
      Polyimides, uses and miscellaneous
     Polyoxyarylenes
Polysulfones, uses and miscellaneous
23
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Urethane polymers, uses and miscellaneous
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (hollow-fiber membranes of, for affinity membrane prepn. for sepn. of
        biomols.)
IT
     Polymers, uses and miscellaneous
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (hollow-fibers membrane of, for affinity membrane prepn. for sepn. of
ΙT
     Bacteria
     Plant cell
        (surface receptor of, conjugates with hollow-fiber membrane
        polymer, for biomol. sepn.)
IT
     Immunoglobulins
     RL: ANST (Analytical study)
        (A, conjugates, antibody to, with hollow-fiber membrane
        polymer, for biomol. sepn.)
ΙT
     Proteins, specific or class
    -RL:-ANST (Analytical study)
        (A, conjugates, with hollow-fiber membrane polymer,
        for biomol. sepn.)
ΙT
     Antigens
     RL: ANST (Analytical study)
        (CEA (carcinoembryonic antigen), antibody to, conjugates with
        hollow-fiber membrane polymer, for biomol. sepn.)
ΙT
     Immunoglobulins
     RL: ANST (Analytical study)
        (E, conjugates, antibody to, with hollow-fiber membrane
        polymer, for biomol. sepn.)
IT
     Immunoglobulins
     RL: ANST (Analytical study)
        (G, conjugatés, antibody to, with hollow-fiber membrane
        polymer, for biomol. sepn.)
ΙT
     Immunoglobulins
     RL: ANST (Analytical study)
        (M, conjugates, antibody to, with hollow-fiber membrane
        polymer, for biomol. sepn.)
ΙT
     Polymerization
        (Ziegler-Natta, hydrophobic polymer for
        hollow-fiber affinity membrane prepn. by)
     Siloxanes and $ilicones, compounds
ΙT
     RL: ANST (Analytical study)
        (arom., conjugates, with hollow-fiber membrane
        polymer, for biomol. sepn.)
ΙT
     Ligands
     RL: ANST (Analytical study)
        (conjugated with hollow-fiber membrane polymer,
        for biomol. sepn.)
ΙT
     Avidins
     Enzymes
     Histones
     Immunoglobulins
     Monosaccharides
    Nucleic acids
     Polysaccharides, compounds
     Siloxanes and Silicones, compounds
     RL: ANST (Analytical study)
        (conjugates suroles ASSIGcalHistonesROLES ASS
        Nucleic acidsous ROL *Polysaccharides, )
    Carbohydrates and Sugars, compounds Glycoproteins, specific or class
IT
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```
Proteins, specific or class
     RL: ANST (Analytical study)
        (conjugates, mbrane polymers, for biomol.
        sepn.Glycoproteins, speci)
     Lymphokines and Cytokines
ΙT
     RL: ANST (Analytical study)
        (interleukins, antibody to, conjugates with hollow-fiber
        membrane polymer, for biomol. sepn.)
ΙT
     Polymerization:
        (ionic, hydrophobic polymer for hollow-fiber
        affinity membrane prepn. by)
ΙT
     Proteins, specific or class
     RL: ANST (Analytical study)
        (ligand-binding, conjugates, with hollow-fiber membrane
        polymer, for biomol. sepn.)
IT
     Antibodies
     RL: ANST (Analytical study)
    -- (monoclonati-conjugates with hollow-fiber membrane -
        polymer, for biomol. sepn.)
IT
     Nucleotides, polymers
     RL: ANST (Analytical study)
        (oligo-, conjugates, with hollow-fiber membrane
        polymers, for biomol. sepn.)
IT
     Nucleotides, polymers
     RL: ANST (Analytical study)
        (poly-, conjugates, with hollow-fiber membrane
        polymers, for biomol. sepn.)
IT
     Vinyl compounds, polymers
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (polymers, halogenated, hollow-fiber membranes of, for
        affinity membrane prepn. for sepn. of biomols.)
IT
     Polymers, compounds
     RL: ANST (Analytical study)
        (polysulfonates, conjugates, with hollow-fiber membrane
        polymer, for biomol. sepn.)
ΙT
     Polymerization
        (radical, hydrophobic polymer for hollow-fiber
        affinity membrane prepn. by)
IT
     Polymerization
        (ring-opening, hydrophobic polymer for hollow-fiber
        affinity membrane prepn. by)
ΙT
     Polymerization
        (stepwise, hydrophobic polymer for hollow-fiber
        affinity membrane prepn. by)
IT
     Dyes
        (synthetic conjugates with hollow-fiber membrane
        polymer, for biomol. sepn.)
ΙT
     Animal growth regulators
     RL: ANST (Analytical study)
        (transforming growth factors, antibody to, conjugates with
        hollow-fiber membrane polymer, for biomol. sepn.)
IT
     Fetoproteins RL: ANST (Analytical study)
        (.alpha.-, conjugates, antibody to, with hollow-fiber
        membrane polymer, for biomol. sepn.)
ΙT
     9002-61-3, Chorionic gonadotropin
                                          9002-71-5, Thyrotropic hormone
     RL: ANST (Analytical study)
        (antibody to, conjugates with hollow-fiber membrane polymer, for biomol. sepn.)
     9004-62-0D, Hydroxyethyl cellulose, linked polysulfone conjugates
ΙT
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```
RL: ANST (Analytical study)
        (for affinity hollow-fiber membrane prepn. for biomol. sepn.)
     58-85-5D, Biotin, polymer-linked conjugates
IT
     9000-11-7D, Carboxymethyl cellulose, polymer-linked
                 9002-89-5D, Poly(vinyl alcohol), polymer
     conjugates
                         9002-98-6D, polymer-linked
     -linked conjugates
                 9004-34-6D, Cellulose, alkyl ethers, linked-
     conjugates
     polymer conjugates 9004-54-0D, Dextran,
     polymer-linked conjugates
                                9005-49-6D, Heparin,
     polymer-linked conjugates
                                9015-73-0D,
     Diethylaminoethyldextran, polymer-linked conjugates
     27357-96-6D, polymer-linked conjugates
     RL: ANST (Analytical study)
        (in affinity hollow-fiber membrane, for biomol. sepn.)
     56-81-5, 1,2,3-Propanetriol, uses and miscellaneous 872-50-4, NMP, uses
IT
     and miscellaneous
     RL: USES (Uses)
        (polyether sulfone/poly(ethylene oxide) polymer dope contg.,
        for affinity hollow-fiber membrane for biomol. sepn.)
     105913-11-9, Plasminogen activator
IT
     RL: ANST (Analytical study)
        (tissue, antibody to, conjugates with hollow-fiber membrane
        polymer, for biomol. sepn.)
L13 ANSWER 14 OF 15 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                         1991:445683 HCAPLUS
DOCUMENT NUMBER:
                         115:45683
                         Process and pulsed alternating voltage enzyme
TITLE:
                         electrode sensor for measuring the glucose content of
                         glucose-containing fluids under anaerobic conditions
                         Kuypers, Martinus Henricus; Steeghs, Gerardus
INVENTOR(S):
                         Fransiscus Jozef
PATENT ASSIGNEE(S):
                         PPG Hellige B. V., Neth.
                         Eur. Pat. Appl., 16 pp.
SOURCE:
                         CODEN: EPXXDW
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     ED 2007
                                           APPLICATION NO. DATE
                     · ----
                                           ----
     EP 396788 A1 19901114
                                           EP 1989-108264
                                                             19890508
         R: AT, CH; DE, ES, FR, GB, GR, IT, LI, NL, SE
     Glucose is measured in liq. media, esp. blood under anaerobic conditions,
AΒ
     using an electrochem. sensor operated with a pulsed alternating voltage
     switchable between a higher operating voltage level (A), at which excess O
     is released at the working electrode and into the surrounding immobilized glucose oxidase by way of electrochem. splitting H2O, and a lower
     operating voltage (B), at which only the catalytic glucose reaction in
     glucose oxidase takes place to form H2O2 which oxidizes at the working electrode. The current flowing thereby is detd. as the value sensed in
     the phase of low operating voltage level B and is evaluated as a measure
     of glucose concn. Other embodiments and diagrams of the sensors are
     given.
IC
     ICM C12M001-40
CC
     9-7 (Biochemical Methods)
IT
     Polymers, uses and miscellaneous
     RL: USES (Uses)
        (glucose oxidase immobilized in, in oxygen-producing pulsed alternating
```

voltage enzyme electrode sensor for glucose detn.) ΙT Electrodes (bio-, enzyme, membrane, hydrophobic, in oxygen-producing pulsed alternating voltage enzyme electrode sensor for glucose detn.) L13 ANSWER 15 OF 115 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1981:84576 HCAPLUS DOCUMENT NUMBER: 94:84576 Ring-opening polymerization of bicyclic oxalactone and TITLE: oxalactam. Speciality polymers having hydrophilic- and hydrophobic microdomains AUTHOR(S): Sumitomo, Hiroshi Fac. Agric., Nagoya Univ., Nagoya, 464, Japan CORPORATE SOURCE: Polym. Prepr., Am. Chem. Soc., Div. Polym. Chem. SOURCE: (1979), 20(1), 134-7CODEN: ACPPAY; ISSN: 0032-3934 DOCUMENT TYPE: Journal LANGUAGE: English A discussion of the prepn. and properties of macrocyclic oligoesters AB (dimers, tetramers, and hexamers) obtained by the ring-opening polymn. of 6,8-dioxabicyclo[3.2.1]octan-7-one (I) or optically active (+)-(1R,5R)-I and of a hydrophilic polyamide membrane obtained from 8-oxa-6azabicyclo[3.211]octan-7-one (II) by simultaneous ring-opening polymn. and casting. 35-3 (Synthetic High Polymers) CC polydioxabicycloctanone oligoester; polyoxaazabicycloctanone membrane; ST polyamide oxaazabicyclooctanone membrane; polyester dioxabicyclooctanone oligomer; cyclic oligomer dioxabicyclooctanone; lactam oxaazabicyclooctanone polymer Membranes and praphragms ΙT (polyamides) contg. alternating amide and tetrahydropyran groups, hydrophilic) ΙT Polyamides, préparation RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of contg. alternating amide and tetrahydropyran

groups, as hydrophilic membranes)

IT 49793-24-0P 76623-37-5P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)

(prepn. and properties of, as hydrophilic membrane)

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=> fil wpids
FILE 'WPIDS' ENTERED AT 10:07:39 ON 07 MAR 2002
COPYRIGHT (C) 2002 DERWENT INFORMATION LTD
FILE LAST UPDATED: 06 MAR 2002
                                            <20020306/UP>
                                      200215
MOST RECENT DERWENT UPDATE
                                               <200215/DW>
DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE
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>>> FOR UP-TO-DATE INFORMATION ABOUT THE DERWENT CHEMISTRY
    RESOURCE, PLEASE VISIT
         http://www.derwent.com/chemistryresource/index.html <<<
>>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES,
    SEE http://www.derwent.com/dwpi/updates/dwpicov/index.html <<<
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     (FILE 'WPIDS' ENTERED AT 10:01:00 ON 07 MAR 2002)
                DEL HIS Y
         109020 S MEMBRANE#
L1
           5262 S (£1)(6A) (ALTER? OR DISRUPT? OR ENHANC? OR STRUCT?)
L2
         443676 S POLYMER##
L3
           5262 S L1 AND L2
L4
         236933 S TRANSPORT?
L5
L6
            226 S E4 AND L5
           1452 S 11 (4A) (ALTER? OR DISRUPT? OR ENHANC? (3A) TRANSPOR?)
L7
            161 S L7 AND L3
L8
              5 S L8 AND ENHANC? (4A) TRANSPOR?
L9
              6 S L8 AND CONJUGAT?
L10
          33823 S HYDROPHOB?
L11
          43851 S HYDROPHIL?
L12
             13 S L8 AND L11 AND L12
L13
             20 S 113 OR L10 OR L9
L14
     FILE 'WPIDS' ENTERED AT 10:07:39 ON 07 MAR 2002
=> d .wp tech 1-20
L14 ANSWER 1 OF 20 WPIDS COPYRIGHT 2002 DERWENT INFORMATION LTD
     2001-602251 [68] WPIDS
DNC C2001-178322 F
     Non-naturally occurring gene therapy vector useful for gene therapy,
ΤI
     comprises an inner shell having a core complex containing a nucleic acid
     and at least one complex forming reagent.
DC
     A96 B04 B05 DÎ6
     CHENG, C; FREI, J; METT, H; PUTHUPPARAMPIL, S; STANEK, J; SUBRAMANIAN, K;
     TITMAS, R; WOODLE, M; YANG, J
     (NOVS) NOVARTIS AG; (NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH
PΑ
CYC
PI
     WO 2001049324 A2 20010712 (200168) * EN 178p
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
            NL OA PT SD SE SL SZ TR TZ UG ZW
         W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM
            DZ EE ESTFI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC
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LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW AU 2001033669 À 20010716 (200169) WO 2001049324 Å2 WO 2000-EP13300 20001228; AU 2001033669 A AU 2001-33669 20001228 AU 2001033669 A Based on WO 200149324 FDT PRAI US 1999-475305 19991230 WO 200149324 A UPAB: 20011121 NOVELTY - A non-naturally occurring gene therapy vector, comprising an inner shell having a core complex (1) containing a nucleic acid and at least one complex forming reagent (2), is new. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following: (1) forming a self assembling core complex by feeding a stream of a solution of a mucleic acid and a core complex-forming moiety into a static mixer, the streams are split into inner and outer helical streams that intersect at several different points causing turbulence and promoting mixing, that frequency in a physicochemical assembly interaction; and (2) a compound having formula (I). m = 3 or 4;Y = -(CH2)n-, or -CH2-CH=CH-CH2- if R2 is -(CH2)3-NR4R5 and m is 3; n = 3-16;R2 = H, or lower alkyl, or -(CH2)3-NR4R5 is m is 3; R3 = H, or alkyl, or -CH2-CH(-X')-OH if R2 is -(CH2)3-NR4R5 and m is 3; X and X independently, H or alkyl; and R, R1, R4 and R5 = independently, H or lower alkyl, where R, R1, R4 and R5 are not all H or methyl, if m is 3 and Y is -(CH2)3. ACTIVITY None given. MECHANISM OF ACTION - Gene therapy. No biological data is given. USE - In gene therapy for nucleic acid delivery. ADVANTAGE: The vectors are stable having an improved outer steric layer that provides enhanced target specificity, in vivo and colloidal stability. The vectors are relatively homogenous and comprises chemically defined species. The vectors demonstrate improved cell entry and intracellular trafficking, permitting enhanced nucleic acid therapeutic activity such as gene expression. Dwg.0/30 UPTX: 20011121 TECH TECHNOLOGY FOCUS - BIOLOGY - Preferred Components: The vector further comprises a fuscgenic moiety, an outer shell moiety and a targeting moiety. The vector comprises a mixture of at least two outershell reagents in which each but the outershell reagents comprises the hydrophilic polymer having substantially different sizes. The fusogenic moiety is incorporated directly in (1) and comprises a shell that is anchored to (1). The fusogenic moiety comprises at least one moiety selected from a viral peptide, an amphiphilic peptide, a fusogenic polymer lipid conjugate and a biodegradable fusogenic polymer-lipid conjugate. The fusogenic moiety is a membrane surfactant polymer-lipid conjugate selected from Thesit (RTM), Brij 58 (RTM), Brij 78 (RTM), Tween 80 (RTM), Tween 20 (RTM), C12E8, C14E8, C16E8, Chol-PEG 900, analog containing polyoxazoline or other hydrophilic polymer substituted for the PEG and analog having fluorogarbons substituted for the hydrocarbon. CnEn = hydrocarbon poly(ethylene glycol) ether; C = hydrocarbon of carbon length N; and E = poly(ethylene glycol) of degree of polymerization N. The inner shell is anchored to the outer shell moiety via a covalent linkage that is degradable by chemical reduction or sulfhydryl treatment

at a pH of at most 6.5. The covalent linkage is selected from  $-C(O)-NH-N=CH+\frac{1}{2}$  -C(O)-NH-NH-C(O)-NH=CH-, -O-T-CH=N-NH-C(O)- or -NH-C(O)-CH2-CH2-S-S-. The outer shell moiety stabilizes the vector and reduces nonspecific binding to proteins and cells. The outer shell moiety is anchored to the fusogenic moiety and (1) and comprises a hydrophilic polymer. The outer shell comprises the targeting molety. The outer shell comprises a protective polymer conjugate in which the polymer exhibits solubility in both polar and non-polar solvents. The targeting moiety enhances binding of the vector to a target tissue and cell population. The targeting element is a receptor ligand, an antibody or antibody fragment, a targeting peptide, a targeting carbohydrate molecule or a lectin, preferably vascular endothelial cell growth factor, fibroblast growth factor (FGF)2, somatostatin and its analog, transferrin, melanotropin, ApoE and ApoE peptide, von Willebrand's Factor and von Willebrand's Factor peptide; adenoviral fiber protein and adenoviral fiber protein peptide; PD1 and PD1 peptide, epidermal growth factor (EGF) and EGF peptide, RGD peptide, folate, pyridoxyl, sialyl-Lewis and chemical analogs. (2) is selected from a lipid, a polymer, and a spermine analog complex of (I). The complex-forming lipid agent is selected from phosphatidylcholine, phosphatidylethanolamine, dioleoylphosphatidylethanolamine, dioleoylphosphatidylcholine, cholesterol and other sterols, N-1-(2,3-dioleyloxy)propyl-N,N,N-trimethylammonium chloride, 1,2 bis (oleoyloxy) -3-(trimethylammonia) propane, phosphatidic acid, phosphatidylglycerol, phosphatidylinositol, glycolipids comprising two optional vunsaturated 14-22C hydrocarbon chains, sphingomyelin, sphingosine, ceramide, terpenes, cholesterol hemisuccinate, cholesterol sulfate, diacylglycerol, 1,2-dioleoyl-3-dimethylammonium propanediol, dioctadecyldimethylammonium bromide, dioctadecyldimethylammonium chloride, dioctadecylamidoglycylspermine, 1,3-dioleoyloxy-2-(6carboxyspermyl propylamide, Lipofectamine7 (RTM) (2,3-dioleyloxy-N-(2-(sperminecarboxamido)ethyl)-N, N-dimethyl-1-propanaminium trifluoroacetate), hexadecyltrimethyl-ammonium bromide, dimethyl-dioctadecylammonium bromide, 1,2-dimyristyloxypropyl-3-dimethylhydroxy ethyl ammonium bromide, dipalmitoylphosphatidylethanolamylspermine , dioctylamineglycinespermine, dihexadecylamine-spermine (C18-2-Sper), aminocholesterol-spermine, 1-(2-(9(Z)-octadecenoyloxy)ethyl)-2(8(Z)heptadecenyl) 3-(2-hydroxyethyl) imidazolinium chloride, dimyristoyl-3-trimethylammonium-propane, 1,2-dimyristoyl-sn-glycero-3-ethylphosphatidylcholine, lysylphosphatidylethanolamine, cholestryl-4 aminoproprionate, Genzyme-67 (spermadine cholestryl carbamate), 2- dipalmitoyl-1,2-propandiol)-4-methylimidazole, 2-(dioleoyl-1,2-propandiol)-4-methylimidazole, 2-(cholestryl-1-propylamine carbamate) imidazole, N-(4-pyridyl)-dipalmitoyl-1,2-propandiol-3-amine, 3-beta-(N-(N',N'- dimethylaminoethane)carbamoyl)cholesterol, 3beta-(N-(N',N(,N),N'-trimethylaminoethane)carbamoyl) cholesterol, 1,2-dioleoyl-sn-glycero-3-succinate, 1,2-dioleoyl-sn-glycero-3-succinyl-2hydroxethyl disulfide ornithine conjugate, 1,2-dioleoyl-snglycero-3-succinyl-2-hydroxethyl hexyl orithine conjugate, N, N', N, N'-tetramethyl-N, N', N, N'-tetrapalmityolspermine, 3-tetradecylamino-N-tert-butyl-N'- tetradecylpropionamidine (vectamidine or diCl4-amidime), YKS-220 (RTM) (N-(3-(2-(1,3-dioleoyloxy)propoxycarbonyl)prog致加N,N,N-trimethyl ammonium iodide) and DC-6-14 (RTM) (O,O'-ditetradecanoyl-N-(alpha-trimethylammonioacetyl)diethanolamine chloride). (2) comprises a mixture of at least two (2). (2) possesses at least one additional activity selected from cell binding, biological membrane fusion endosome disruption and nuclear targeting. The nucleic acid is selected from a recombinant plasmid, a replication-deficient plasmid, a mini-plasmid, a recombinant viral genome, a linear nucleic acid fragment, an antisense agent, a linear

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polynucleotide, a circular polynucleotide, a ribozyme, a cellular promoter
and a viral genome. (2) further comprises a nuclear targeting moiety that
enhances nuclear binding and/or uptake. The nuclear targeting moiety is
selected from a nuclear localization signal peptide, a nuclear membrane
transport peptide or a steroid receptor binding moiety. The nuclear
targeting modely is anchored to the nucleic acid in (1). The viral peptide
is selected from MLV env peptide, HA env peptide, a viral envelope protein
ectodomain, a membrane-destabilizing peptide of a viral envelope protein
membrane-proximal domain, a hydrophobic domain peptide segment
of a viral fusion protein or an amphiphilic-region containing peptide. The
amphiphilic-region containing peptide is selected from melittin,
magainins, fusion segments from Haemophillus influenza hemagglutinin (HA)
protein, human immunodeficiency virus (HIV) segment I from the cytoplasmic
tail of HIV 1gp41 or amphiphilic segments from viral env membrane
proteins.
TECHNOLOGY FOCÜS - POLYMERS - Preferred Components: The
fusogenic moiety comprises a fusogenic polymer, a fusogenic
polymer lipid conjugate, a biodegradable fusogenic
polymer or a biodegradable fusogenic polymer-lipid
conjugate. (2) is a polymer of structure
-(-N(R1)-CH2-R2-)x-(-N(R3)-CH2-R2-)y-. The fusogenic moiety is a
polymer of structure - (-N(R1)-CH2-R2-)x-(-N(R'3)-CH2-R2-)y-.
R1 and R3 = hydrocarbon optionally substituted with amine, guanidinium
or imidazole moiety;
R2 = lower alkyl;
x and y = not defined;
R'3 = hydrocarbon optionally substituted with carboxyl, hydroxyl,
sulfate or phosphate.
The outer shell comprises a protective steric polymer
conjugate in which the polymer is selected from the
group consisting of polyethylene-glycol (PEG), a polyacetal
polymer, a polyocazoline polymer optionally block with
end-group conjugation, a hydrolyzed dextran polyacetal
polymer, a polyoxazoline, a polyethylene glycol, a
polyvinylpyrrolidone, polylactic acid, polyglycolic acid,
polymethacrylamide, polyethyloxazoline, polymethyloxazoline,
polydimethylagtylamide, polyvinylinethylether, polyhydroxypropyl
methacrylate, polyhydroxypropylmethacrylamide, polyhydroxyethyl acrylate, polyhydroxyethyloxazoline, polyhydroxypropyloxazoline, polyaspartamide or
a polyvinyl alcohol.
ANSWER 2 OF 20 WPIDS COPYRIGHT 2002
                                         DERWENT INFORMATION LTD
2001-596316 [67]
                    WPIDS
C2001-176449
Composition, for disruption of cell membrane, used for
delivering dīāģnostic or therapeutic agents to cytoplasm of cells,
contains hydrophobic polymer and hydrophilic
component coupled via linkage which is cleaved as function of pH.
HOFFMAN, A S; MURTHY, N; STAYTON, P S
(UNIW) UNIV WASHINGTON
93
WO 2001051092 A2 20010719 (200167)* EN
                                           50p
   RW: AT BE CHICY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
       NL OA PT SD SE SL SZ TR TZ UG ZW
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       DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC
       LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU ZA ZW
AU 2001027648 A 20010724 (200168)
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WO 2001051092 A2 WO 2001-US356 20010105; AU 2001027648 A AU 2001-27648 ADT 20010105 FDT AU 2001027648 A Based on WO 200151092 PRAI US 2000-174893P 20000107 WO 200151092 A UPAB: 20011119 NOVELTY - Composition, for disruption of membrane, contains conjugate comprising: (1) Polymer which is hydrophobic under conditions where membrane is to be disrupted; and (2) Hydrophilic component which is an agent to be delivered, or groups/polymer linkable/linked to hydrophobic polymer, in amount making conjugate hydrophilic. The hydrophilic component is coupled to hydrophobic polymer via linkage which is cleaved as function of pH. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for a method of producing the composition and a method of disrupting a cell or organelle using the composition. USE - The composition is used for delivering diagnostic or therapeutic agénts, through cell membranes; barriers; or layers, to cytoplasm of cells, and for the release of cell contents for subsequent recovery and/or analysis. ADVANTAGE - The composition can deliver agents to cells without significant lysosomal degradation, and can be controlled externally by non-invasive means such as ultrasound. Dwq.0/6 UPTX: 20011119 TECH TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Components: The hydrophilic component is a therapeutic, diagnostic, or prophylactic agent to be delivered to a cell or organelle, preferably a protein, peptide, nucleotide, saccharide, polysaccharide, preferably a nucleotide molecule selected from antisense, ribozyme, ribozyme guide sequence, triplex forming nucleotide and gene, complexed to a polymer component of the conjugate. The hydrophobic polymer is vinyl-type, non-vinyl, or naturally derived. The hydrophilic group is a hydroxy acid, thiol, amine, carboxyl, or amino acid. The conjugate further comprises a ligand specifically binding to a target molecule and the composition further comprises a carrier selected from carriers for systemic, local, or topical delivery of the conjugate.
Preferred Linkages: The hydrophilic groups are coupled directly to the hydrophobic polymer. The linkage coupling the hydrophilic to the membrane disruptive component is dissruptable upon exposure to physical or chemical stimulus, it is preferably stable at pH 6.8 - 8, disrupted at pH less than 6.5, hydrolyzes within 30 - 60 minutes at pH 5.0 - 5.5, and is acetal, orthoester, cis-aconityl, carboxylic acid hydrazone, phosphamide, ester, Schiff base, vinyl ether, dithioacetal, tert butyl ester, carbamate, urethane, annyaride, polysaccharide, amide, ester, ether, thiourea, urea, thioester, sulfenamide, phosphoroamidate, or amine N-oxide. The agent to be delivered is coupled to the hydrophilic or membrane disruptive component by a degradable or dissruptable linkage, Preferred Cell The cell is in a patient, an endosome in a cell, or a bacterial cell

ANSWER 3 OF 20 WPIDS COPYRIGHT 2002 L14 DERWENT INFORMATION LTD

2001-582012 [65] AN WPIDS

C2001-172537 DNC

Compositions and methods for enhancing drug delivery across biological

membranes, using a delivery-enhancing
transporter having guanidino or amidino moieties.

DC B05

IN ROTHBARD, J B; WENDER, P A

PA (CELL-N) CELLGATE INC; (ROTH-I) ROTHBARD J B; (WEND-I) WENDER P A

CYC 94

PI WO 2001062297 A1 20010830 (200165) \* EN 54p

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001036920 A 20010903 (200202) US 2002009491 A1 20020124 (200210)

ADT WO 2001062297 AT WO 2001-US4459 20010209; AU 2001036920 A AU 2001-36920 20010209; US 2002009491 Al Provisional US 2000-182166P 20000214, US 2001-779693 20010207

FDT AU 2001036920 A Based on WO 200162297

PRAI US 2001-779693 20010207; US 2000-182166P 20000214

AB WO 200162297 A UPAB: 20011108

NOVELTY - Drug delivery across biological membranes and tissues, particularly across 1 or more layers of skin, is enhanced using delivery-enhancing transporters having guanidino or amidino moieties.

DETAILED DESCRIPTION - A method for delivery of a compound to the surface of, into or across a biological barrier, comprises contacting the barrier with a composition comprising the compound and a delivery enhancing transporter comprising sufficient guanidino or amidino moieties to increase delivery of the compound compared to delivery in the absence of the transporter. An INDEPENDENT CLAIM is included for a composition comprising a biologically active agent and a delivery enhancing transporter comprising guanidino or amidino moieties.

ACTIVITY Antiinflammatory; antiulcer; antiallergic; antiasthmatic; vasotropic; antiparkinsonian; neuroleptic; cytostatic; anti-HIV; anticonvulsant; neuroprotective; tranquilizer; vulnerary; antidepressant; nootropic; antimigraine; analgesic.

MECHANISM OF ACTION - H2 histamine inhibitor; proton-potassium ATPase inhibitor.

The ability of polyArg to facilitate cellular uptake of small organic acids was determined. In separate vials, n equivalents (n = 1-6) of fluorescein (poorly soluble in water) were added to the free base of a nonamer of arginine in water. Phosphoric acid (6-n equivalents) was added to each flask, and the solutions were frozen and lyophilized. When the dried powders were taken up in water, they were very water soluble. The 8 compounds had fluorescein:peptide ratio from 1:1 to 6:1.

When dilutions of each of the solutions were used in cellular uptake

When dilutions of each of the solutions were used in cellular uptake assays, the resultant cells were stained equivalently, showing that all fluorescein molecules were deposited on the cell surface. The staining pattern of the cells was different when compared to fluorescein that was covalently attached to short polymers of arginine. Distinct punctate staining was seen on the cell surface as well as in the cytosol, when covalent conjugates were used. Staining of individual cells was very heterogeneous, with the variation in cell fluorescence ranging over 3 orders of magnitude. However, when noncovalent conjugates were used, cell staining was uniform with cell fluorescence varying only by a factor of 2-4. Staining was intense, with the majority of the dye on the cell surface.

USE - For delivery of drugs and diagnostic imaging or contrast

agents, across biological membranes and tissues, e.g. cell membranes, mitochondrial membranes, dermal and epithelial membranes, and across the blood-brain barrier. The compound may be delivered into and across the stratum corneum, stratum ganulosum, stratum lucidum and/or stratum germinativum.

The compositions can be used to treat e.g. Crohn's disease, ulcerative colitis, gastrointestinal ulcers, peptic ulcer disease or abnormal proliferative disease; a bronchial condition (e.g. cystic fibrosis, asthma, allergic rhinitis and chronic obstructive pulmonary disease); ischemia, Parkinson's disease, schizophrenia, cancer, acquired immune deficiency syndrome, infections of the central nervous system, epilepsy, multiple sclerosis, neurodegenerative disease, trauma, depression, Alzheimer's disease, migraine, pain or a seizure disorder. Dwg.0/5

TECH

UPTX: 20011108

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Drugs: The drug is an antiviral agent, e.g. acyclovir, famciclovir, ganciclovir, foscarnet, idoxuridine, sorivudine, trifluridine, valacyclovir, cidofovir, didanosine, stavudine, zalcitabine, zidovudine, ribavirin or rimantatine; an antibacterial agent, e.g. nafcillin, oxacillin, penicillin, amoxicillin, ampicillin, cefotaxime, ceftriaxone, rifampin, minocycline, ciprofloxacin, norfloxacin, erythromycin or vancomycin; an antifungal agent, e.g. amphotericin, itraconazole, ketoconazole, miconazole, nystatin, clotrimazole, fluconazole, ciclopirox, econazole, naftifine, terbinafine or griseofulvin; an antineoplastic agent, e.g. pentostatin, 6-mercaptopurine, 6-thioguanine, methotrexate, bleomycins, etoposide, teniposide, dactinomycin, daunorubicin, doxorubicin, mitoxantrone, hydroxyurea, stiluorouracil, cytarabine, fludarabine, mitomycin, cisplatin, procedure, dacarbazine, paclitaxel, colchicine or vinka alkaloids; immunosuppressive agents, e.g. methotrexate, azathioprine, fluorouracil, hydroxyurea, 6-thioguanine, cyclophosphamide, mechloroethamine hydrochloride, carmustine, cyclosporine, taxol, tacrolimus, vinblastine, dapsone or sulfasalazine; an analgesic agent, e.g. lidocaine, bupivacaine, novocaine, procaine, tetracaine, benzocaine, cocaine, mepivacaine, etidocaine, proparacaine, ropivacaine or prilocaine; a vitamin; or hormone. Preferred Transporter: The delivery enhancing transporter is preferably a peptide having 6-15 amino açı́aβresidues, where 6-12 are selected from L-arginine, D-arginine, L-homoarginine or D homoarginine. Preferred Method: The compound is an H2 histamine inhibitor, an inhibitor of the proton-potassium ATPase or an antibiotic directed at Helicobacter pylori. The compound is delivered into and across stratum corneum, stratum granulosum, stratum lucidum or stratum germinativum. The compound is a diagnostic or contrast agent.

- ANSWER 4 OF 20 WPIDS COPYRIGHT 2002 L14DERWENT INFORMATION LTD
- 2001-091577 [10] AN WPIDS
- DNC C2001-027033
- Electrochemical sensor for subcutaneous implantation into a mammal's body TIto measure an analyte in subcutaneous fluid has working electrode and analyte responsive sensing layer contacting with the analyte only at an edge of the sensor.
- DC B04 D16
- AUDETT, J D; CHO, B; SAKSLUND, H; SAY, J; TOMASCO, M F; YAMASAKI, D IN
- PΑ (THER-N) THERASENSE INC
- CYC 92
- WO 2000078992 A2 20001228 (200110) \* EN PI 42p
  - RW: AT BE CHRCY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW
    W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ

EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SE TJ TM TR TT TZ UA UG UZ VN YU ZA ZW

AU 2000057471 Å 20010109 (200122)

WO 2000078992 A2 WO 2000-US16773 20000616; AU 2000057471 A AU 2000-57471 ADT 20000616

AU 2000057471 A Based on WO 200078992 FDT

PRAI US 2000-194618P 20000405; US 1999-139936P 19990618

WO 200078992 A UPAB: 20010220

NOVELTY - An electrochemical sensor (100), comprising a working electrode (104) and an analyte-responsive sensing layer (134) near the working electrode, is hew. The sensing layer is exposed to contact with the analyte only at an edge of the sensor. The sensor signal is limited, at least in part, by mass transport of an analyte to the sensing layer.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for a method of determining the concentration of an analyte in a body fluid of a mammal, comprising:

- (a) implanting at least a portion of the novel electrochemical sensor into the body of a mammal so that the edge of the sensor contacts body fluid of the mammal; and
- (b) measuring the concentration of the analyte in the body fluid using the sensor.
- USE For subcutaneous implantation into the body of a mammal for contact with body fluids of the mammal to measure an analyte in subcutaneous fluid (claimed).

ADVANTAGE → The invention restricts the mass transport of the analyte to the sensing layer eliminating the need for a mass transport limiting membrane. The enhancement of operational stability and operational life of the sensor includes:

- (a) reduced flux of analyte to the sensing layer which reduces the rate of enzyme turnover;
- (b) deeper diffusion of glucose into the sensing layer to reach relatively unused enzyme as enzyme at the edge of the sensing layer is deactivated during use;
- (c) the immobilization of the sensing layer between the base and top layers which limits the swelling of a hydrogel; and
- (d) the reduction of the risk of the enzyme leaching into tissue. DESCRIPTION OF DRAWING(S) - The figure shows a perspective view of an analyte sensor

Electrochémical sensor 100

Sensor body 101

Working electrode 104

Reference/counter electrode 108

Top surface 111

Base layer 112

Base layer 113

Top layer 116

Proximal end 120

Distal edge 124

Side edge 128

Analyte-responsive sensing layer 134.

Dwg.1/16

UPTX: 20010220 TECH

TECHNOLOGY FOCUS - INSTRUMENTATION AND TESTING - Preferred Device: The sensor has a flexible body (101) having an inner peripheral surface extending into the sensor. The edge, is a peripheral edge, side edge (128), or preferably a distal edge (124) of the sensor, which is planar or preferably cylindrical. T edge at which the sensing layer is exposed is defined by at least a portion of the inner peripheral surface. The sensor has a base layer (113) and an oxygen permeable top layer (116), both

layers being impervious to analyte, and a less than 100, preferably 1-10 micro-m thick spacer layer. The sensing layer is non-leachably disposed on the sensor. The sensing layer and the spacer layer are at least partially disposed between these base and top layers. The spacer layer defines a channel with apperipheral surface extending into the spacer layer, and the edge at which the sensing layer is exposed is defined by at least a portion of this inner peripheral surface.

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Materials: The analyte is glucose, and the sensing layer comprises a redox polymer, an enzyme, and a cross-linker.

L14 ANSWER 5 OF 20 WPIDS COPYRIGHT 2002 DERWENT INFORMATION LTD 2001-031893 [04] AN WPIDS DNC C2001-009771 TI Hydrophilic charged microporous membrane useful as filter device for removing bacterial contaminants from water, saline solution, comprises porous hydrophobic substrate and coating of charge-providing agent. DC D15 J01 ISHEE, M; KINSEY, J L; KONSTANTIN, P; SHERTOK, J; WU, X; YANG, Y IN PA (PALL) PALL CORP CYC WO 2000069549 A1 20001123 (200104)\* EN PΙ 42p RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SP SE SL SZ TZ UG ZW W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW AU 2000050036 2 20001205 (200113) ADT WO 2000069549 A1 WO 2000-US12894 20000512; AU 2000050036 A AU 2000-50036 20000512 FDT AU 2000050036 A Based on WO 200069549 PRAI US 1999-134197P 19990514 WO 200069549 A UPAB: 20010118 NOVELTY - A hydrophilic charged microporous membrane comprises a porous hydrophobic substrate and a coating comprising charge-providing agent.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following: (i) preparing hydrophilic charged microporous membrane by contacting porous hydrophobic substrate with composition containing charge-providing agent or its precursor; (ii) device comprising hydrophilic charged membrane; and (iii) process for treating fluid containing bacterial contaminants by placing fluid in contact with the hydrophilic charged microporous

membrane and récovering a bacterial contaminant depleted fluid.

USE - As filter device (claimed) for separation, or removal of bacterial contaminants from water, saline solution and other fluids. For filtering biological fluid such as lymph and cerebrospinal fluid and pharmaceutical products such as composition containing proteins (e.g. antibodies, enzyme, vaccines), amino acids, peptides, nucleic acids, plasmids, cosmids, phages, polysaccharides, lipids, bioreactor, fermenter and/or cell culture harvests.

ADVANTAGE. The membrane has effective endotoxin retaining capacity, water and/or saline solution wettability and water permeability.

Dwg.0/7

TECH UPTX: 20010118

TECHNOLOGY FOCUS - POLYMERS - Preferred Component: The hydrophobic substrate or matrix is a polymer such as

polyethersulfone substrate or matrix. The charge-providing agent is distributed within hydrophobic polymer matrix. The porous hydrophobic substrate (substantially) free of wetting agent. Preferred Agent: The charge-providing agent is positively charged or negatively charged. The positively charged agent is positively charged polymer (PCP). The PCP contains quarternary ammonium groups. The PCP is polyamine (or acrylic polymer) containing quarternary ammonium groups. The polyamine is crosslinked through ring opened epoxy groups. The acrylic polymer comprising polymerized acrylol monomer preferably alkacryloyl monomer, more preferably alkacryloylaminoalkyl monomer is crosslinked through a polyfunctional crosslinking agent (such as alkylene glycol diacrylate). A negatively charged polymers comprises sulfonic acid groups, preferably polymerized acrylamido sulfonic acid monomer. The negatively charged polymer is cross linked by acrylamidione cross linking agent (such as N-(alkoxymethyl) acrylamide. The negatively charged polymer further includes polymerized hydroxyalkylarcylate monomer or polyacrylate such as ethylene glycol or dimethyl acrylate . Preferred Polymer : The polymer is poly aromatics, polysulfones, polyolefins, polystyrenes, polyamides, polyimides, fluoropolymers, polycarbonate, polyesters on cellulose acetate. Preferred Process: In the coating process the substrate is contacted with composition containing charge-providing agent and cured and the membrane is extracted to remove the residue. Alternately the hydrophilic charged microporous membrane is obtained by forming a casting solution containing polymer capable of forming porous hydrophobic matrix, solvent for polymer, pore former, and charge-providing agent or its precursor. The casting solution is shaped to obtain a pre-membrane by causing phase-inversion to obtain a phase-inverted membrane. The phase-inverted membrane is leached. Preferred Precursor: The precursor comprises a free radical polymerizable monomer, a crosslinking agent and free radical initiator. The free radical polymerizable monomer is positively charged acrylic monomer containing quarternary ammonium group or negatively charged free radical polymerizable monomer containing sulfonic acid group. The cross linking agent is polyacrylate preferably diacrylate such as alkylene glycol diacrylate.

TECHNOLOGY FOCUS - ENVIRONMENT - Preferred Contaminated Fluid: The fluid contaminated with bacteria is water or pharmaceutical product such as saline solution with surface tension of 72-78 dynes/cm.

TECHNOLOGY FOCUS - BIOLOGY - Preferred Contaminants: The bacterial contaminant comprises an endotoxin.

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ANSWER 6 OF 20 WPIDS COPYRIGHT 2002
L14
                                                 DERWENT INFORMATION LTD
     2000-389642 [34] WPIDS
AN
     2000-423196 [36]
CR
DNC
     C2000-118513
TI
     Compositions comprising membrane- and micelle forming lipids for
     delivering pharmaceuticals to organisms.
DC
     A96 B02 B07
ΙN
     LEIGH, S
      (PHAR-N) PHARES PHARM RES NV
PΑ
CYC
                   Å, 20000614 (200034)*
ΡI
     GB 2344520
                                                   13p
ADT GB 2344520 A GB 1998-27006 19981208
PRAI GB 1998-27006 19981208
AB GB 2344520 A UPAB: 20000801
     NOVELTY - A composition (I) for delivering a biologically active compound
     to an organism comprising a pharmaceutically active compound dissolved or
```

dispersed in a lipid and a **polymer** for modifying the swelling properties of the composition and/or for rendering the composition comminutable or friable, is new.

ACTIVITY T Variable

MECHANISM OF ACTION - Pharmaceutical carrier.

No data given.

USE - The composition (I) is used for delivering pharmaceutically active agents, such as nifedipine and griesofulvin (claimed), to a patient.

ADVANTAGE - (I) has improved physiological properties and loading capacities allowing the development of novel dosage forms. (I) provides maximum bioavailability with minimum side effects. It is an improved carrier for both lipophilic and hydrophobic active compounds that is safe, effective and may provide benefits in a range of applications.

In particular, (I) provides:

- (1) enhanced ability to effect molecular solution of poorly water soluble compounds;
- (2) enhanced absorption of both hydrophilic and lipophilic compounds through lipid-membrane interactions and altered permeability; and
- (3) longer in vivo retention of the hydrated lipid associate on absorption surfaces.

Dwg.0/0

TECH UPTX: 20000718

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Composition: (I) may be in comminuted form, spheroidized form, pellet form or in the form of a tablet or capsule.

(I) preferably comprises at least 1 therapeutically active compound and at least 1 micelle forming lipid. The compound is at least partly in suspension on the lipid (preferably a mixture of membrane lipids and micelle-forming lipids).

The therapeutically active compound is nifedipine or griesofulvin depending on the mix of membrane lipids and micelle-forming lipids utilized.

TECHNOLOGY FOCUS - POLYMERS - Preferred Composition: The polymer comprises 1-50% by weight (wt%) of the composition. The polymer is a methacrylic resin, a povidine, a cellulose derivative, a polyvinyl alcohol and/or polyvinyl phthalate or is a gum. preferably, it is an acrylic polymer whose degree of swelling depends on the pH.

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Composition: The lipid is a membrane lipid and/or a micelle forming lipid. They are monoacyl lipids and diacyl lipids present in the weight ratio of 1:99 to 99:1 (preferably 1:10 to 10:1)

The lipid comprises 1-20 wt% (especially 10 wt%) of the composition. Preparation: The monoacyl and diacyl lipids are a mixture obtained by enzyme hydrolysis.

- L14 ANSWER 7 OF 20 WPIDS COPYRIGHT 2002 DERWENT INFORMATION LTD
- AN 2000-389588 [34] WPIDS
- DNN N2000-291751 DNC C2000-118496
- TI Specific immunoblotting test for protein antigens, used particularly to detect erythropoietin in sportsmen, with transfer of bound primary antibody to second membrane before detection.
- DC A96 B04 D16 J04 S03
- IN LASNE, F
- PA (HOSP-N) HOSPICES CIVILS LYON ETAB

31p

CYC

PI FR 2786273 A1 20000526 (200034)\*

ADT FR 2786273 A1 FR 1998-14864 19981120

PRAI FR 1998-14864

19981120

AB FR 2786273 A UPAB: 20000718

NOVELTY - A qualitative and/or quantitative immunoblotting assay of one or more target protein antigens (Ag) in which complexes formed on a first membrane are dissociated and the released primary antibodies (Ab1) are transferred to a transfer membrane (Mt) where they are detected by reaction with secondary antibody (Ab2) is new.

DETAILED DESCRIPTION - A qualitative and/or quantitative immunoblotting assay of one or more target protein antigens (Ag) in which complexes formed on a first membrane are dissociated and the released primary antibodies (Ab1) are transferred to a transfer membrane (Mt) where they are detected by reaction with secondary antibody (Ab2) is new. The test sample is applied to a membrane (Mdb) directly or Ag are separated (from each other and form other proteins) on at least one separation support (Ss) and transferred to an absorption membrane (Mwb) to form an image of Ss. Mab or Mwb is saturated, preferably with a solution of inert protein, then reacted with one or more specific Ab1 to form complexes. The membranes are then washed to remove unbound Ab1. The bonds in any Ag/Ab1 complexes on the membrane are then broken by altering the physico-chemical conditions and/or the environment in and on the membrane and released Abl are transferred, particularly by desorption, to Mt, leaving Ag and other proteins on the first membrane, therefore forming an image of Abl on Mt. The transferred Abl are then reacted with Ab2, Mt washed to remove unbound Ab2 and any formation of Ab1/Ab2 complexes detected. Optionally the information is captured on at least one other support, particularly by exposure of a sensitive film.

INDEPENDENT CLAIMS are also included for the following:

- (1) device for transfer of released Ab1 to Mt; and
- (2) kit for performing the assay including the device of (a).
- USE The method is particularly used to detect (recombinant) erythropoietin or other illicit drugs in humans or animals, especially to detect drug abuse by sportsmen, but may also be used to detect cellular, serum or viral proteins.

serum or viral proteins.

ADVANTAGE Transfer of Abl to a separate membrane overcomes the problem of false positives associated with non-specific reaction of Ab2 with other proteins on the membrane, since an Ag/Abl/Ab2 complex is never formed on the first membrane. The method is generally applicable (to any antigen of Ab2), provides high detection sensitivity for targeted Ag and is simple and economical to perform.

Dwg.0/5

TECH

UPTX: 20000718

TECHNOLOGY FOCUS - BIOLOGY - Preferred conditions: The initial complex is disrupted by reducing the pH to 4 or lower, particularly 1-5 (sic) for a temperature of 15-25 degreesC. The disruption (and subsequent) steps are performed as soon as primary complexes are detected on the first membrane. Preferably before transfer of released Ab1, the first membrane and Mt are placed face-to-face, under pressure, and Mt is saturated before treatment with Ab2.

Preferred Materials: All membranes are of natural or synthetic polymers. Ab2 are labeled with a chemiluminescent system, e.g. with biotin for subsequent reaction with streptavidin/peroxidase conjugate and then chemiluminescent substrate. Ag are particularly (recombinant) erythropoietin (EPO) or cellular, serum or viral proteins, or their mixtures. Agents for saturating membranes are e.g. skim milk or serum albuming and suitable test samples, optionally diluted, are blood, urine, saliva, cerebro-spinal fluid, cell culture media and intracellular fluids.

Preferred Process: Where Ag are separated initially, this is by electrophoresis or isoelectric focusing.

TECHNOLOGY FOCUS - INSTRUMENTATION AND TESTING - Preferred Device: The device of (1) consists of two plates that retain, in sandwich fashion, the first and second membranes in face-to-face contact. Preferably it also includes an arrangement for wetting the membranes, particularly a piece of absorbent paper soaked in liquid medium.

TECHNOLOGY FOCUS - POLYMERS - Suitable membrane materials are halogenated polyalkylidenes (e.g. poly(vinylidene fluoride)) or optionally modified cellulose (e.g. nitrocellulose and/or cellulose acetate).

ANSWER 8 OF 20 WPIDS COPYRIGHT 2002 L14 DERWENT INFORMATION LTD

1999-633730 [54] WPIDS AN

DNC C1999-185055

New conjugate of lipid with basic, membrane-TΙ disrupting peptide having reversed amide backbone, used to introduce animatic macromolecules or active agents into cells, e.g. for gene therapy.

DC B04 D16

IN KITAS, E A; SCHLAEGER, E

PA (HOFF) ROCHE DIAGNOSTICS GMBH

CYC 84.

PΙ WO 9951629 A2 19991014 (199954)\* EN 23p

RW: AT BE CONCY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SP SE SL SZ UG ZW

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE GH GM HR HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MX MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VŅ YU ZA ZW

AU 9937065 Å 19991025 (200011)

A2 20010117 (200105) EN EP 1068225

R: AT BE CH DE DK ES FI FR GB GR IE IT LI LU NL PT SE WO 9951629 A2 WO 1999-EP2361 19990407; AU 9937065 A AU 1999-37065

19990407; EP 1068225 A2 EP 1999-919208 19990407, WO 1999-EP2361 19990407 FDT AU 9937065 A Based on WO 9951629; EP 1068225 A2 Based on WO 9951629 PRAI EP 1998-124837 19981230; EP 1998-106302 19980407

9951629 A UPAB: 19991221

NOVELTY - Conjugates of (i) lipid and (ii) basic,

membrane-disrupting peptides, and their salts, are new.

DETAILED DESCRIPTION - Conjugates of (i) lipids and (ii)

basic, membrane disrupting peptides of formula (I) and their salts, comprise:

R1 and R2 = residues of linear or branched, saturated or unsaturated aliphatic carboxylic acids or phospholipids;

R3 = basic, membrane-disrupting peptide with a reversed amide backbone;

Y = 2-100 alkylene;

= CONH or SS

INDEPENDENT CLAIMS are also included for the following:

- (a) the pertide QQRKRKIWSILAPLGTTLVKLVAGIC-NH2 (II) with a reversed amide backbone and with at least 50% of residues D-amino acids, and its
- (b) compositions containing (I), at least one of helper lipid, short-chain phospholipid and/or cationic lipid, optionally also an additional transfection reagent; and
- (c) process for introducing into a cell, in vivo or in vitro, an anionic macromolecule (III) or a biologically active anionic molecule (IIIa), by treating the cells with (III) or (IIIa) in presence of (I). ACTIVITY None given.

MECHANISM OF ACTION - None given.

USE - (I) are used to introduce, into eukaryotic or prokaryotic cells, in vivo or in vitro, anionic macromolecules, particularly nucleic acids, or biologically active anionic molecules. In particular DNA (for gene therapy or recombinant protein production), antisense sequences, haptamers, triplex-formers, ribozymes etc. also proteins and peptides (for immunization) are introduced.

ADVANTAGE - Transfection with (I) provides rapid expression of heterologous proteins in large scale systems, with both adherent and suspended cells, even at low DNA concentrations and without significant inhibition by conditioned medium. A conjugate prepared from 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine N-(3-(2pyridyldithio) propionate and the all D reversed-backbone peptide OORKRKIWSILAPLGTTLVKLVAGIC-NH2 was formulated with a plasmid encoding the human tumor necrosis receptor protein p55 and used to transfect HEK293 (EBNA) cells. At a conjugate concentration of 10 mg/ml, cell viability was 90-95% with p55 expression 83 ng/ml. Dwg.0/0

UPTX: 19991221 TECH

> TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Compounds: In (I), R1 and R2 are acyl residues of a 12-20C carboxylic acids, especially lauroyl, palmitoyl, stearoyl or oleoyl; X = disulfide; R3 = R3'-CH(CONH2)-CH2-; R3' = residue of (II) without terminal Cys.

Preferred peptide: (II) has all amino acids, except Gly, in D-configuration.

Preparation: Either;

(i) R3NH2 is reacted with a carboxy-lipid (i.e. (I) with XR3 replaced by carboxy); or

(ii) R3SH is reacted with a derivatized lipid, i.e. (I) with XR3 replaced with -SZ, where Z is a leaving group such as 2-pyridylthio. The lipid derivatives are known from Biochim. Biophys. Acta, 862 (1986) 435, and the peptides are produced by standard methods of solid-phase

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Compositions: These also include (III), particularly a polynucleotide, and optionally also a polycationic polymer, particularly poly(ethylene imine). They are formulated as aqueous or organic solutions or dispersions, or as a liposome or micelle. Helper lipids are e.g. phosphatidyl ethanolamines and short phospholipids are dicapryl- or dicaproyl-phosphatidyl choline. The optimal mole ratio of (I):helper lipid is 1-10; of helper lipid:short phospholipid 2:20 and of (I) and additional transfection component 0.1:10.

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Method: For transfection with nucleic acid, the ratio of positive charges to negative charges between (I) and (III) is 0.1-10, preferably 0.5-5, typically using 0.1-10, especially 0.2-2, mg of (III) per 104 cells, in vitro, or doses of 0.0001-1 g in vivo.

ANSWER 9 OF 20 WPIDS COPYRIGHT 2002 L14 DERWENT INFORMATION LTD

AN

1999-540689 [45] WPIDS N1999-400749 DNC C DNN DNC C1999-157934

TΙ Ion conductive matrixes for forming membranes, composite electrode, electrochemical cell, fuel cell and water electrolizer. A32 A85 E16 E36 E37 J03 L03 P56 X16

DÇ

IN

DUVDEVANI, T; MELMAN, A; PELED, E (UYRA-N) UNIV RAMOT APPLIED RES & IND DEV LTD PΑ

CYC

PΙ WO 9944245 Å1 19990902 (199945)\* EN 35p

RW: AT BE CHOCY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG ZW

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG UŚ UZ VN YU ZW

Å 19990915 (200004) Å1 20010110 (200103) AU 9926369 EP 1066656

R: DE ES FR GB IT NL SE

A 20001206 (200103) IL 123419

IL 126830 A. 20010520 (200153)

KR 2001034536 A 20010425 (200164)

WO 9944245 Alimo 1999-IL109 19990222; AU 9926369 A AU 1999-26369 19990222; EP 1066656 AllEP 1999-906424 19990222, WO 1999-IL109 19990222; IL 123419 A IL 1998-123419 19980224; IL 126830 A IL 1998-126830 19981030; KR ·2001034536 A KR 2000-709294 20000823

FDT AU 9926369 A Based on WO 9944245; EP 1066656 Al Based on WO 9944245 PRAI IL 1998-126830 19981030; IL 1998-123419 19980224

9944245 A UPAB: 19991103

NOVELTY - The ion conductive matrix comprises 5 - 60 volume percent. (vol.%) of inorganic powder in form of sub-micron particles having good aqueous electrolyte absorption capacity, 5 - 50 vol.% of polymeric binder compatible with an aqueous electrolyte, and 10 - 90 vol.% of an aqueous electrolyte.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for: (i) Method for casting membrane which comprises preparing mixture comprising inorganic powder, polymeric binder, at least one high boiling point solvent with boiling point above 100 deg. C and at least one low boiling point solvent in which the polymeric binder is soluble or forms a gel at casting temperature. Film is casted out of mixture and low boiling point solvent is evaporated from mixture to form solid film. Solid film is washed to replace high boiling point solvent with aqueous electrolyte solution. Alternatively, mixture is heated to its softening température and film is formed by hot extrusion of softened mixture. The high boiling point solvent used in the mixture has boiling point above 90 deg. C. Film is cooled to obtain solid film, and washed to replace solvent with aqueous electrolyte solution.

(ii) Method for casting composite electrode comprising steps involved in casting membrane. Alternatively, preparing composite electrode by extrusion which comprises steps involved in

preparing membrane by extrusion.

USE - For forming membranes, composite electrode, electrochemical cell, fuel cell, and water electrolizer.

ADVANTAGE Novel, low cost and highly conductive ion conducting matrix, membranes and electrodes are provided. The ion conducting membranes have good porosity and mechanical properties. Internal lubricants with low solubility in water is used to achieve solubility factor not higher than 14 (cal/cc)1/2, thereby preventing the migration of internal lubricants out of ion conductive membranes when they come in contact with water at washing phase or acid loading phase. Dwq.0/2

UPTX: 19991103 TECH

> TECHNOLOGY FOCUS - INORGANIC CHEMISTRY - Preferred Matrix: The ion conducting matrix is a proton conducting matrix and comprises desirably 5 - 50 % of inorganic powder such as silicon dioxide (SiO2), zirconium oxide (ZrO2), boron trioxide (B2O3), titanium oxide (TiO2), aluminum oxide (Al2O3), and or optional hydroxides or oxy-hydroxides of Ti, Al, B or Zr with a surface area of at least 10 m2/g. The matrix optionally comprises 0.1 - 25 % of nonvolatile liquid lubricant which is compatible with all the components in matrix.

> Preferred Electrolyte: The aqueous electrolyte consists of aqueous soluble salt and/or base which is used in aqueous solution having molar

concentration of 0.1 - 10 M, preferably 1 - 5 M. Alkali metal salts, alkali earth metal salts, R4NX, where R = organic radical; X = anion derived from an inorganic acid. Ammonium chloride (NH4Cl) and/or zinc chloride (ZnCl2) is used as the aqueous soluble salt. R4NOH, where R = hydrogen or an organic radical, alkali and/or alkali earth base compound is used as the aqueous soluble bases. Preferred Membrane: The membrane comprises ion conducting matrix having electronically nonconductive inorganic material with particle size less than 150 nm. The membrane comprises pores with size less than 50 nm. The inorganic powder of matrix is treated with acid or base prior to preparation of membrane. The membrane further comprises electronic nonconductive reinforcing element. Preferred Electrode: The composite electrode comprises 10 - 70 vol.% of the matrix and remaining electrode material. Preferred Electrochemical Cell: The electrochemical cell comprises membrane or at least one electrode having electrode material of carbon and/or graphite, metal oxides such as RuO2, WOx or MnO2. Cadmium, zinc, and/or aluminum or its alloys is used as anode active material. Manganese oxide (MnO2) # silver oxide or nickel oxy hydroxide (NiOOH) is used as cathode active material. Zn or Al anode and oxygen or air electrode which consists of double layer film with hydrophobic air side and hydrophilic fonic membrane side is used. The air electrode catalyst is compatible with aqueous solutions of ionic conductive membrane such as oxides of platinum, palladium, gold, silver, copper, manganese, tungsten and for metal-porphyrin complexes of their salts. The electrochemicalicell is single structure unit manufactured by hot pressing the electrodes on both sides of the membrane.

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Lubricant: Diesters of aliphatic or aromatic dibasic acids, esters of phosphoric acids, hydrocarbons or synthetic hydrocarbons, silicone oils and/or fluorocarbons is used as the lubricant.

Preferred Acid: The proton conducting matrix comprises 10 - 90 vol.% of an acid such as CF3(CF2)nSO3H, HO3S(CF2)nSO3H, where n = 0 - 9, especially 0 - 4.

sulfuric acid, hydrochloric acid, hydrobromic acid, phosphoric acid and/or nitric acid. The acid is used in an aqueous solution having a molar concentration of 10 - 99 %, preferably 25 - 99 %.

Preferred Solvent: The high boiling point solvent used for casting or preparing memorane or composite electrode is water soluble solvent.

Propylene carbonate, ethylene carbonate, dimethyl phthalate, diethyl phthalate, and/or dibutyl phthalate is used as high boiling point solvent for casting or preparing membrane. Tetrahydrofuran, dimethylether (DME), cyclopentanone, acetone, N-methyl pyrrolidone, dimethylacetamide, methylethylketone, and/or dimethyl-formamide is used as the low boiling point solvent for casting or preparing the membrane. Propylene carbonate, diethyl carbonate, dimethyl carbonate, butyrolactone, methyl isoamyl ketone, cyclonexanone, dialkyl phthalate, and/or glycerol triacetate is used as solvent for casting or preparing composite electrode.

TECHNOLOGY FOCUS - POLYMERS - Preferred Binder: Polyvinylidene fluoride, polyvinylidene fluoridehexafluoropropylene, poly(tetrafluoroethylene), poly(methylmethacrylate), polysulfone amide, poly(acrylamide), polyvinyl chloride, poly(acrylonitrile), and/or polyvinyl fluoride is used as the polymeric binder.

L14 ANSWER 10 OF 20 WPIDS COPYRIGHT 2002 DERWENT INFORMATION LTD

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1999-468807 [39]
                          WPIDS
ΑN
DNC
     C1999-137478
     Composition for enhancing transport through cell
ΤI
     membranes, particularly for delivery of genes or toxins -
     comprises transporting agent such as polymer that changes
     structure or properties in response to stimulus..
     A14 A96 A97 B04 B07 D16
DC
     CRUM, L A; HOFFMAN, A S; LACKEY, C; MOURAD, P D; MURTHY, N; PORTER, T M;
IN
     PRESS, O; STAYTON, P; TIRRELL, D; PRESS, O W
     (UYMA-N) UNIV MASSACHUSETTS; (UNIW) UNIV WASHINGTON; (CRUM-I) CRUM L A;
PΑ
     (HOFF-I) HOFFMAN A S; (LACK-I) LACKEY C; (MOUR-I) MOURAD P D; (MURT-I)
     MURTHY N; (PORT=1) PORTER T M; (PRES-I) PRESS O W; (STAY-I) STAYTON P;
     (TIRR-I) TIRRELL D
CYC
    23
                    A1 19990715 (199939)* EN
PΙ
     WO 9934831
                                                  52p
        RW: AT BE CHOCY DE DK ES FI FR GB GR IE IT LU MC NL PT SE
         W: AU CA JP
     AU 9920261 19990726 (199952)
EP 1044021 11 20001018 (200053) EN
         R: AT BEACH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE
     US 2001007666 A1 20010712 (200143)
     JP 2002500201 W 20020108 (200206)
                                                  62p
     wo 9934831 Alimo 1999-US122 19990105; AU 9920261 A AU 1999-20261 19990105;
     EP 1044021 AT EP 1999-900750 19990105, WO 1999-US122 19990105; US
     2001007666 Al Provisional US 1998-70411P 19980105, US 1999-226044
     19990105; JP 2002500201 W WO 1999-US122 19990105, JP 2000-527278 19990105
FDT AU 9920261 A Based on WO 9934831; EP 1044021 A1 Based on WO 9934831; JP
     2002500201 W Based on WO 9934831
PRAI US 1998-704118 19980105; US 1999-226044
                                                     19990105
           9934831 A UPAB: 19990928
AΒ
     WO
     NOVELTY - Composition for enhancing transport, or
     release, through cell membranes, between cells, cell barriers or
     lipid membranes comprises:
           (1) a membrane barrier transporting agent (I) and
           (2) system for inducing, or enhancing, effectiveness of (I) for
     membrane disruption.
          DETAILED DESCRIPTION - (I) is:
           (1) a polymer (Ia) that changes structure or properties in
     response to some stimulus;
           (2) a hydrophobic peptide (Ib) that forms pores in cell membranes as
     a function of thange in pH, or
           (3) a phospholipid disrupting agent (Ic).
          ACTIVITY None given.

MECHANISM OF ACTION - Disruption of cell membranes
          USE - The composition is particularly used to improve delivery, to
     cells, of diagnostic or therapeutic agents, including nucleic acids,
     proteins, synthetic compounds, metals, radiolabels etc., particularly for
     gene therapy, ê.g. treatment or prevention of restenosis, or toxins such
     as ricin for killing of target cells. It may also be used to release
     metabolites or other analytes from cells, for subsequent measurement.

ADVANTAGE The compositions can be controlled and manipulated
     externally, by noninvasive methods.
           DESCRIPTION OF DRAWING(S) - Graph showing inhibition of cellular
     protein synthesis by ricin toxin A chain (RTA); poly(propylacrylic acid), PPAAc, and a mixture of the two, showing increased delivery of RTA in presence of the polymer.
     Dwg.6/7
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TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Materials: System (b) may

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induce a change in structure of (Ia) and is then an alteration in pH,
    light, ionic strength, solvent composition, temperature or electric field.
    Alternatively, (b) comprises ultrasound, an electric field and/or
    radiation to provide an enhancing effect, particularly ultrasound of
     frequency 20 kHz to 10 MHz for delivery through the skin. Preferred
    Composition: This may also include:
     (1) a diagnostic or therapeutic agent (especially a cytotoxic compound,
    nucleoside, nucleotide or nucleic acid, ionically or covalently linked to
     (I)), particularly where (Ia) is pH sensitive, i.e. responsive at pH
     5-6.5, the conditions present in an endosome;
     (2) an agent that decreases lysosomal degradation, e.g. an enzyme
     inhibitor or vėrapamil;
     (3) a polycationic polymer, e.g. polylysine or chitosan;
     (4) a carrier leg. micro- or nano-particles, liposomes, emulsions or
     lipid vesiclés;
     (5) an agent that increases endocytosis, e.g. an antibody (which may also
     serve as targeting agent).
    Process: The composition is applied to a suspension of cells, to layers of
     cells, or to lipid membranes, optionally in conjunction with
     electrophoresis or iontophoresis.
     TECHNOLOGY FOCUS - POLYMERS - pH-sensitive polymers
     are graft, block or random polymers of acrylic acid or its 1-6C
     linear, branched or cyclic 2-alpha-alkyl derivatives, acrylate
     ester/acrylic acid copolymers, or polymers containing at least
     one block of protein or peptide that includes imidazole groups.
    ANSWER 11 OF 20 WPIDS COPYRIGHT 2002
                                             DERWENT INFORMATION LTD
     1999-045281 [04]
                      WPIDS
    C1999-014190
    Enhancing transport across biological membrane
     - comprises containing membrane with conjugate containing active
     agent covalently attached to transport polymer.
     ROTHBARD, J B; WENDER, P A
     (STRD) UNIV LELAND STANFORD JUNIOR
    83
                  A2 19981126 (199904) * EN
                                              50p
    WO 9852614
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL
            OA PT SD SE SZ UG ZW
        W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE GH GM GW HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG
           MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG
           US UZ VN YU ZW
    AU 9875938
                  À 19981211 (199917)
                 Å2 20000202 (200011)
                                        ĖΝ
     EP 975370
        R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
            RO SE SI
                 Å 20000315 (200016)
Å3 20000517 (200031)
     GB 2341390
     CZ 9904066
    ADT
     19980521; EP \frac{197}{5}370 A2 EP 1998-923711 19980521, WO 1998-US10571 19980521;
     GB 2341390 A WO 1998-US10571 19980521, GB 1999-23841 19991011; CZ 9904066
     A3 WO 1998-US10571 19980521, CZ 1999-4066 19980521; CN 1263473 A CN
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1998-807186 ½9980521; GB 2341390 B WO 1998-US10571 19980521, GB 1999-23841
     19991011; AU 734827 B AU 1998-75938 19980521; KR 2001012809 A KR
     1999-710772 19991120; BR 9809138 A BR 1998-9138 19980521, WO 1998-US10571
     19980521; US 6306993 B1 Provisional US 1997-47345P 19970521, US 1998-83259
     19980521; JP 2002502376 W JP 1998-550716 19980521, WO 1998-US10571
     19980521
    AU 9875938 A Based on WO 9852614; EP 975370 A2 Based on WO 9852614; GB
     2341390 A Based on WO 9852614; CZ 9904066 A3 Based on WO 9852614; GB
     2341390 B Based on WO 9852614; AU 734827 B Previous Publ. AU 9875938,
     Based on WO 9852614; BR 9809138 A Based on WO 9852614; JP 2002502376 W
     Based on WO 9852614
PRAI US 1997-47345P 19970521; US 1998-83259
                                                 19980521
          9852614 A UPAB: 20001117
    WO
      Enhancing the transport of a selected compound across
     a biological membrane comprises contacting the membrane with a
     conjugate containing a biologically active agent covalently
     attached to a transport polymer so that the contacting promotes
     the transport of the conjugate across the membrane at a greater
     rate than the transport of the non-conjugated biological agent
     across the membrane. The polymer comprises 6-25 subunits, at
     least 50% of which contain a guanidino or amidino side chain group and the
    polymer contains at least 6 continuous guanidino and/or amidino
     side chain groups. Also claimed are a composition for delivering a
     biologically active agent across a biological membrane and an excipient
     and a combinatorial library of conjugates.
          The biologically active agent preferably includes small organic
     compounds e.g. the anticancer taxane, antimicrobial agents (against
     bacteria or fungi such as yeast), polypeptides e.g. protein antigens,
     proteins, oligosaccharides, nucleic acids and metal ions. The agent has a
    molecular weight of < 10 kDa. The agent may be linked to the
    polymer via a linker group, preferably a cleavable linker e.g. a
     linker group that is cleavable by an enzyme or by solvent-mediated
     cleavage such as an ester, amide or disulphide group, or it may contain a
     photocleavable group.
          USE - The method is used to transport biologically active agents
     across biological membranes including eukaryotic and prokaryotic cell
     membranes. The method may be used to screen conjugates for
     biological activity. The conjugates are formed form candidate
     agents including candidate agents selected from a combinatorial library.
     Dwg.0/7
    ANSWER 12 OF 20 WPIDS COPYRIGHT 2002
                                             DERWENT INFORMATION LTD
     1998-362918 [3]]
                        WPIDS
DNN
    N1998-283296
                        DNC C1998-111762
     Glucose sensor for use in low-oxygen environments, useful e.g. in diabetes
     - comprises an enzyme-containing membrane made from a semi-
     interpenetrating polymer network which increases oxygen
     transport to the enzyme.
     A96 B04 D16 J04 S03
     BLUBAUGH, E A BRUNSMAN, A R
     (IMPL-N) IMPLANTED BIOSYSTEMS INC
CYC
     23
                  A1 19980625 (199831) * EN
                                              29p
     WO 9827419
        RW: AT BE CHODE DK ES FI FR GB GR IE IT LU MC NL PT SE
        W: AU CA JP SI
                 A 19980715 (199846)
     AU 9856116
                  A1 19991013 (199947)
                                        EN
     EP 948743
         R: AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE
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A 19991012 (199949) B 20000608 (200035)

US 5964993 AU 720712

ΑN

TΙ

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AB

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JP 2001506365 20010515 (200133)
ADT WO 9827419 A1 0 1997-US23426 19971217; AU 9856116 A AU 1998-56116
     19971217; EP 348743 A1 EP 1997-952530 19971217, WO 1997-US23426 19971217;
     US 5964993 A 05 1996-769863 19961219; AU 720712 B AU 1998-56116 19971217;
     JP 2001506365職 WO 1997-US23426 19971217, JP 1998-527981 19971217
FDT AU 9856116 A Based on WO 9827419; EP 948743 Al Based on WO 9827419; AU
     720712 B Previous Publ. AU 9856116, Based on WO 9827419; JP 2001506365 W
     Based on WO 9827419
PRAI US 1996-769863 19961219
          9827419 A UPAB: 19991122
     An enzyme-containing membrane comprises a semi-interpenetrating
    polymer network of fibrillated polytetrafluoroethylene and a
     silicon compound, in which the network is infiltrated with an enzyme. Also
     claimed are: (1) a membrane system useful in a sensor in which efficient
     oxygen transport is desired, comprising an outer membrane and an inner
     membrane which is an enzyme-containing membrane as above, and (2) a
     glucose sensor comprising: a membrane system with an inner and outer
     membrane, in which the inner membrane is an enzyme-containing membrane as
     above, and the outer membrane restricts the flow of glucose into the inner
     membrane; an electrode capable of oxidising hydrogen peroxide; and in
     which the inner membrane is in between the outer membrane and the
     electrode.
          The silicon compound is a cross-linked polyorganosiloxane, especially
     a polydimethylsiloxane. The enzyme is capable of oxidising glucose and
     generating hydrogen peroxide. It is an oxidase, preferably glucose
     oxidase, and is immobilised within the network as an enzyme gel. The
     membrane has porosity of 25-55%, and contains 15-40 vol.% silicon
     compound. The buter membrane comprises polycarbonate. The inner membrane
          USE - The products are used for measuring glucose levels in
     low-oxygen enwironments, in an implantable device, especially useful in
     diabetes.
          ADVANTAGE? The sensor accurately measures glucose levels, even in
     low-oxygen environments e.g. biological fluids. Oxygen is needed in the
     reaction involved in the measurement process, and the inner
     membrane enhances the transport of oxygen to
     the site of glucose oxidation.
     Dwq.0/3
    ANSWER 13 OF 20 WPIDS COPYRIGHT 2002
                                             DERWENT INFORMATION LTD
T.14
     1990-329157 [44]
                        WPIDS
ΑN
DNC C1990-142890
TI
     Prepn. of microporous membrane - from mixt. of hydrophobic
     polymer and hydrophilic polymer comprising
     leaching part of hydrophilic polymer from matrix.
DC
     A14 A26 A88 J01
     KOENHEN, D MANULDER, M H; ROESINK, H D W; SMOLDERS, C A; MULDER, M H V
IN
PA
     (XFLO-N) X-FLOW BV
CYC
    17
                   Å 19901031 (199044)*
PΙ
     EP 395133
         R: AT BE CA DE ES FR GB GR IT LI LU NL SE
                 A 19901116 (199049)
A 19901028 (199104)
     NL 8901090
     CA 2015413
                  19901214 (199105)
     JP 02302449
                  Å 19911231 (199204)
B1 19950201 (199509)
     US 5076925
     EP 395133
                                         EN
         R: AT BE CH DE DK ES FR GB GR IT LI LU NL SE
     DE 69016491 (E 19950316 (199516)
                 T3 19950601 (199528)
19970225 (199720)
     ES 2070263
     CA 2015413
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$\begin{aligned} \begin{aligned} \begin{aligned} \begin{aligned} \begin{aligned} \begin{aligned} 200147 \end{aligned} \end{aligned} \]
     JP 3196029
                                                    4p
    EP 395133 A EP 1990-200879 19900410; NL 8901090 A NL 1989-1090 19890428;
ADT
     JP 02302449 A:JP 1990-111580 19900426; US 5076925 A US 1990-510070
     19900417; EP 395133 B1 EP 1990-200879 19900410; DE 69016491 E DE
     1990-616491 19900410, EP 1990-200879 19900410; ES 2070263 T3 EP
     1990-200879 19900410; CA 2015413 C CA 1990-2015413 19900425; JP 3196029 B2
     JP 1990-111580 19900426
     DE 69016491 E Based on EP 395133; ES 2070263 T3 Based on EP 395133; JP
FDT
     3196029 B2 Previous Publ. JP 02302449
PRAI NL 1989-1090
                      . 19890428
            395133 A UPAB: 19930928
AB
     A process for preparing a more or less hydrophillic micro porous
     membrane comprises, i) dissolving a hydrophobic polymer
     and a hydrophillic polymer in a suitable solvent or
     mix of solvents, ii) coagulating in a coagulation bath, iii) removing the
     so obtained membrane from the coagulation bath, (iv) and subsequently
     leaching at least a part of the hydrophilic polymer
     from the matrix v) alternatively followed by
     hydrophobisation. The membrane formed thus may be in a
     flat or tubular form or in the form of hollow fibres. Also claimed is a
     microporous membrane comprising essentially of a hydrophobic
     polymer and a more or less hydrophillic polymer
     , which the latter has been cross-linked and has been fixated in or at the
     polymer matrix. The membrane has pores of 0.0001-5 micron, a heat
     resistance of up to 250 deg.C a water permeability of up to 8000 1/mz L
     bar. It also has good chemical resistance and mechanical strength.
           USE/ADVANTAGE - The membranes prepd. are suitable for membrane
     separations, based on particle sizes e.g. ultra-and microfilterations.
     They can also be used as aeration medium, oxygenerator, bioreactor etc.
     0/0
     ANSWER 14 OF 20 WPIDS COPYRIGHT 2002
                                                 DERWENT INFORMATION LTD
     1990-218847 [29]
                          WPIDS
AN
DNN
     N1990-169835
                          DNC C1990-094502
     Modifying surface properties of substrates partic. contact lens - by
ΤI
     (hydro)peroxidising using ozone to direct graft copolymerisation. on
     surface.
DC
     A14 A35 A96 D22 P42 P81
     FREEMAN, E M; JANSSEN, R A; MCCRAW, E C
IN
PΑ
     (CIBA) CIBA GEIGY AG
CYC
     23
                   Å 19900718 (199029)*
PΙ
     EP 378511
          R: AT BE OH DE ES FR GB GR IT LI LU NL SE
                   A 19900719 (199037)
A 19900806 (199037)
     AU 9047936
     NO 9000098
                   A 19900713 (199039)
A 19900714 (199040)
A 19900731 (199041)
     CA 2007552
     FI 9000136
     PT 92815
                   A 19900911 (199042)
     JP 02228309
                   Å 19901106 (199047)
Å 19930818 (199340)
     US 4968532
     IL 92983
                   j Bl 19941102 (199442)
     EP 378511
                                            ΕN
          R: AT BE CH DE DK ES FR GB GR IT LI LU NL SE
                   E 19941208 (199503)
T3 19950201 (199511)
B 19951009 (199545)
     DE 69013698
     ES 2064699
     NO 178072
                   B 19951101 (199604)
B 19970415 (199721)
     IE 65586
     FI 98631
     EP 378511 A EP 1990-810005 19900104; JP 02228309 A JP 1990-1633 19900110;
     US 4968532 A 📆 1989-297018 19890113; IL 92983 A IL 1990-92983 19900105;
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EP 378511 B1 EP 1990-810005 19900104; DE 69013698 E DE 1990-613698
     19900104, EP 1990-810005 19900104; ES 2064699 T3 EP 1990-810005 19900104;
     NO 178072 B NO 1990-98 19900109; IE 65586 B IE 1990-137 19900112; FI 98631
     B FI 1990-136 19900110
    DE 69013698 E Based on EP 378511; ES 2064699 T3 Based on EP 378511; NO
     178072 B Previous Publ. NO 9000098; FI 98631 B Previous Publ. FI 9000136
PRAI US 1989-297018 19890113
           378511 A UPAB: 19930928
    EΡ
     Modifying the surface characteristics of a preformed polymer
     substrate to impart hydrophilicity, hydrophobicity or
     other desired properties by graft polymn. on the substrate
     having (hydro) peroxy gps. on the polymer, of an ethylenically
     unsatd. monome\dot{\hat{\mathbf{r}}}_{i} is claimed. The improvement comprises carrying out the
     graft polymn. on the substrate which is swollen with a liq.
     before or after ozonation, the monomer being insoluble in the liq.
     prevent penetration of the monomer into the interior of the substrate and
     to direct graft polymn. of the monomer to the substrate surface.
          Pref._the process is carried out in the presence of a variable metal
     ion to suppress homoplymsn. of the monomer during grafting, partic a
     ferrous ion, and the ozonation is carried out in water, air, oxygen, or a
     perhalogenated hydrocarbon medium (PHM) (pref.).
          Also claimed is a modification process where the polymer
     substrate is contacted with a soln. which is or contains a chain transfer
     agent (CTA) to saturate or swell the polymer, the soln. being
     insoluble in the PHM, then ozonated and the monomer graft polymn
     . on the substrate surface.
          USE/ADVANTAGE - The process can be used to modify a contact lens
     (claimed) or other biomedical device semipermeable membrane, film or
     fibre, which is modified in respect to hydrophilicity, hydro
     phobicity, opical-, transmission-or bacterial-properties dyeability or
     tintability, opacity, diffraction differences, wettability, bonding
     characteristics, oxygen permeability, lubricity and multilayer
     membrane technology (claimed). Desired alteration of the
     surface characteristics is achieved while maintaining substrate structural
     integrity. @
     0/0
    ANSWER 15 OF 20 WPIDS COPYRIGHT 2002
     ANSWER 15 5 [19]
                                              DERWENT INFORMATION LTD
L14
                        WPIDS
AN
                        DNC C1990-064685
DNN
    N1990-114484
     Membrane for iontophoretic agent delivery device - in which prevents
ΤI
     passive release of drug with release of drug controlled by electric
     current.
     A96 B07 P33 P34 S05
DC
     GYORY, J R; HAAK, R P; THEEUWES, F
IN
PΑ
     (ALZA) ALZA ÇÕŘÉ
CYC
                  À 19900419 (199019)*
PΙ
     WO 9003825
        RW: AT BE CH DE FR GB LU NL SE
         W: AU DK FT IT JP KR NO US
     PT 91890
                      19900430 (199022)
                 A 19900501 (199029)
A 19910402 (199127)
     AU 8944254
     FI 9101589
                      19910717 (199129)
     EP 436658
         R: AT BE CH DE FR GB IT LI LU NL SE
                  19910616 (199129)#
19910603 (199135)
     ES 2019517
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12p

12p

NO 9101186

DK 9100594 US 5080646

US .5147296

A 19910403 (199135) A 19920114 (199206) A 19920915 (199240)

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w 19921015 (199248)
                                               15p
     JP 04505861
     US 5169382
                  Α
                     19921208 (199252)
                                               12p
                      19921208 (199252)
                                               14p
     US 5169383
                  . À
     AU 9229876
                   . A
                      19930211 (199313)
                  À 19930803 (199332)
À 19940621 (199424)
                                               12p
     US 5232438
     US 5322502
                                               12p
                  B 19950406 (199522)
     AU 658246
                   .¢ 19950822 (199540)
     CA 1336781
                   C 19951010 (199548)
     CA 1337300
                  B1 19961014 (199928)
     KR 9614099
     EP 931564
                  A1 19990728 (199934)
         R: AT BE CH DE FR GB IT LI LU NL SE
     JP 11216192
                   À 19990810 (199942)
                                               17p

⇒B1 20000229 (200018)

     US 5232438
     EP 436658 A EP 1989-911931 19891002; US 5080646 A US 1988-252402 19881003;
     US 5147296 A Div ex US 1988-252402 19881003, US 1991-751276 19910828; JP
     04505861 W JP 1989-511045 19881002, WO 1989-US4318 19881002; US 5169382 A
     Cont. of US 1988 = 252402 19881003, US 1991 - 648269 19910130; US 5169383 A WO
     1989-US4318 19891002, US 1990-571577 19900907; AU 9229876 A AU 1992-29876
     19921203, Div ex AU 1989-44254
                                             ; US 5232438 A Cont of US
     1988-252402 19881003, Cont of US 1991-648269 19910130, US 1992-898618
     19920615; US 5322502 A Cont of US 1988-252402 19881003, Cont of US
     1991-648269 tỷ9910130, Cont of US 1992-898618 19920615, US 1993-3761
     19930113; AU 658246 B AU 1992-29876 19921203, Div ex AU 1989-44254
     ; CA 1336781 CDiv ex CA 1989-614338 19890928, CA 1994-616939 19941024; CA
     1337300 C CA 1989-614338 19890928; KR 9614099 B1 WO 1989-US4318 19891002,
     KR 1990-701164 19900602; EP 931564 Al Div ex EP 1989-911931 19891002, EP
     1999-201332 19891002; JP 11216192 A Div ex JP 1989-511045 19891002, JP
     1998-326831 19891002; US 5232438 B1 Cont of US 1988-252402 19881003, Cont
     of US 1991-648269 19910130, US 1992-898618 19920615
    US 5147296 A Div ex US 5080646; JP 04505861 W Based on WO 9003825; US
     5169383 A Based on WO 9003825; US 5232438 A Cont of US 5080646, Cont of US
     5169382; US 5322502 A Cont of US 5080646, Cont of US 5169382, Cont of US
     5232438; AU 658246 B Previous Publ. AU 9229876; EP 931564 Al Div ex EP
     436658; US 5232438 B1 Cont of US 5080646, Cont of US 5169382
PRAI US 1988-252402 19881003; ES 1989-4346
                                                  19891222
          9003825 A UPAB: 19930928
AB
     Membrane for controlling agent delivery from an iontophoretic agent
     delivery device adapted to deliver the agent through an intact body
     surface is claimed, the device having a reservoir contg. the agent to be
     delivered and being connectable to a source of electrical power for
     driving the agent from the reservoir and through the body surface, in
     which the membrane is interposed between the agent reservoir and the body
     surface, the membrane permitting electrically-assisted flux (JEK) of the
     agent and preventing passive flux (Jp) of the agent, the membrane
     exhibiting a (JEK + Jp)/Jp ratio of at least 4, a voltage drop across the
     membrane of less than 1 volt and a Jp of less than 100 microgram/hr-cm2.
          The membrane may be formed by dissolving in a solvent, eg CH2C12 and
     CH3OH, 60-95 pts. wt. of cellulose acetate and 5-40 pts. wt. of a water
     soluble material, eg. polyethylene glycol, having a mol. wt. at least
     great as the agent mol. wt., casting the membrane, evapg. the solvent and
     leaching out all of the water soluble material. Alternatively
     the membrane may comprise a mixt. of a hydrophilic
     resin, eg. PVP or an ion exchange resin having a functional gp. selected
     from sulphonicacid, carboxylic acid, imidodiatcetic acid and quaternary
     amines and hydrophobic polymer, eg. an ethylene vinyl
     acetate polymer having a vinyl acetate content of 1-40 wt. Also claimed is a method for testing performance characteristics of an
     iontophoretic agent delivery device adapted for delivering an agent
     through an intact body surface, using the membrane.
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ADVANTAGE - Using the membrane, the passivee release of drug from the device is prevented and the release of drug is controlled by the magnitude of the electric current. Even if the skin is compromise the amt. of drug delivered ise controlled to a safe level. The membrane can have passive and electrically-assisted transport characteristics similar to that of skin and can be used to test performance characteristics iontophoretic del ANSWER 16 OF 20 WPIDS COPYRIGHT 2002 DERWENT INFORMATION LTD 1988-292714 [41] WPIDS DNC C1988-129791 DNN N1988-222169 Air treatment device contg. volatile active ingredient - has block of erodable gelicast on membrane with liq partially penetrating membrane during casting; A26 A88 A96 B07 C03 D22 P14 P34 SANTINI, T F (DLAI-N) DE LAÌRE INC; (DELA-N) DELAIRE INC CYC 17 WO 8807383 - #A 19881006 (198841) \* EN 51p RW: AT BE CH DE FR GB IT LU NL SE W: AU BR DK A 19880913 (198849) ZA 8802046 A 19881102 (198904) AU 8815736 A 19890307 (198912) US 4809912 16p À 19890405 (198914) EP 309549 R: BE CH DE FR GB IT LI NL (19890330 (198916) PT 87095 WO 8807383 A WO 1988-US1014 19880324; ZA 8802046 A ZA 1988-2046 19880322; US 4809912 A US 1987-32047 19870327; EP 309549 A EP 1988-903665 19880324 PRAI US 1987-32047 19870327 8807383 A UPAB: 19930923 WO An air treatment device has a block of erodible gel enclosed in a vessel whose mouth is closed by a porous membrane. The gel is cast onto the

membrane and liquid partially penetrates the membrane so that the solidified generally attached to the membrane which will support the weight of the cast gel.

The membrane is pref. made from hydrophillic cellulosic fibres coated with polysiloxane or polydimethylsiloxane. Alternatively the membrane is a non-woven synthetic

textile bonded by a water retarding binder.

🖗 🛕 19880615 (198824)

USE/ADVÁNTAGE - The gel has a volatile active ingredient e.g. a fragrance, pheromone, hormone, insecticide, insect attractant, pharmaceutical agent of verterinary drug. The gel is attached to the membrane.

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ANSWER 17 OF 20 WPIDS COPYRIGHT 2002
                                               DERWENT INFORMATION LTD
L14
     1987-348651 [49]
N1987-261223
AN
                         WPIDS
DNN
                         DNC C1987-148972
     Membrane having controlled capillarity - promoting effective contact
TI
     between cells and the membrane surface, used esp. in a blood-typing
     device.
DC
     A96 D16 J04 SQ3
     HEWETT, G
IN
     (GENE-N) GENELABS INC
PΑ
CYC
PΙ
                      19871203 (198749) * EN
     WO 8707304
                                                32p
        RW: AT BE EF DE FR GB IT LU NL SE
         W: AU JP KR
                   A 19871222 (198813)
     AU 8773957
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EP 270569

R: AT BE CH DE FR GB IT LI LU NL SE

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JP 01500369 W 19890209 (198912)
                   A 19890725 (198937)
                                                  qe
     US 4851210
    WO 8707304 A W@ 1987-US838 19870414; EP 270569 A EP 1987-903106 19870414;
     JP 01500369 W P 1987-502892 19870414; US 4851210 A US 1986-866350
     19860522
PRAI US 1986-866350 19860522
          8707304 A UPAB: 19930922
     A structure having controlled capillarity for use in contacting cells in
     soln. with a membrane comprises (a) a porous membrane surface and (b) a
     porous interior capable of incorporating the soln. at a controlled rate
     sufficient to effectively contact the cells with the membrane surface.
     Pref. the membrane is composed of polymer fibres composed of
     e.g. PVDF, PTFE, modified nylon, nitrocellulose, regenerated cellulose or
     cellulose.
          Absorbent porous materials are modified to produce a material having
     controlled capillarity through a 2-step process. The first step is to
     select a base membrane that exhibits a moderately-hydrophilic
     character. The second step is the coating of the membrane with chemical
     agents that control the capillary action of the base membrane materials.
     One example of such coating agents is polymers which make the
     base membrane slightly more hydrophobic. The coating soln.
     contg. e.g. polymer, salt and surfactant is applied after the
     affinity ligand e.g. antibody, has been bound to the membrane.
          USE/ADVANTAGE - The structure has a water-permeable, non-cell
     disruptive membrane that wicks the aqs. phase into a
     hydrophilic matrix thereby efficiently contacting exposed
     antibodies with cells. The structure is used esp. in a blood typing system
     and eliminates the possibility of technical or electrical errors. The
     binding reaction is easily read visibly.
     2/5
     ANSWER 18 OF 20 WPIDS COPYRIGHT 2002
                                                DERWENT INFORMATION LTD
L14
     1987-265616 [38] WPIDS C1987-112499
AN
DNC
     Cast microporous membrane - formed from a soln. of a pore-forming
TI
     polymer and a film-forming polymer and opt. a salt.
DC
     A18 A28 A32 A96 D16 J01
     KETRARO, R; LINDER, C; PERRY, M
IN
PΑ
     (MEMB-N) MEMBRÂNE PROD KIRYA
CYC
                   A 19870923 (198738)* EN
                                                 12p
PΙ
     EP 238276
         R: CH DE FR GB IT LI NL
    JP 63165111 A 19880708 (198833)
US 4761233 A 19880802 (198833) 11p
EP 238276 A EP 1987-302205 19870316; JP 63165111 A JP 1987-62313 19870317;
ADT
US 4761233 A US 1987-24327 19870310
PRAI IL 1986-78169 19860317
AB EP 238276 A UPAB: 19930922
     Prepn. of a microporous membrane comprises casting the membrane from a
     soln. comprising (a) a mixt. of at least one pre-forming polymer
     (I) and at least one film-forming polymer (II), (b) a solvent
     for the mixt of polymers and opt. (c) at least one salt, where (I) is one which if cast alone would contract to form either large pores
     or a non-uniform distribution of material.
           Suitable (I) are e.g. halomethylated polyphenyleneoxide, polystyrene
     derivs., nitrated polysulphones, polyisoprenes and halogenated
     polysulphones Suitable (II) are polysulphones, vinylidene fluoride
     polymers, PTFE based copolymers and polyacry-lonitrile. Suitable
     salts are LiHco3, LiCl, MgCl2, MgCl04, ZnCl2, ZnBr2 and ZnI2.
```

side of the membrane to allow higher fluxes. The membranes are

hydrophilic balance or are readily charged to increase hydrophobicity without redn. in flux. Biologically active

modified easily to alter the hydrophobic/

USE/ADVANTAGE - The membranes have well defined surface pore shapes and the pores are larger on the surface and decrease in size to the bottom

components may be crosslinked to prevent dissolution in organic solvents and to minimise copaction under high pressure. Biologically active members can be used as reactors e.g. enzyme membrane reactors or in chromatographic sepn., or in affinity chromatography for removing specific biological species from complex mists. /3 ANSWER 19 OF 20 | WPIDS COPYRIGHT 2002 DERWENT INFORMATION LTD L141986-070217 [11] WPIDS AN DNC C1986-029943 Sepg. fluid mixt. - by locally heated pervious membrane. ΤI DC J01 ..... ini i IN CHMIEL, H PA (FRAU) FRAUNHÖFER-GES FORD ANGE CYC 13 A 19860306 (198611)\* 13p PΙ DE 3518871 A 19860313 (198612) DE WO 8601425 RW: AT BE CH DE FR GB IT LI NL SE W: JP US A 19860910 (198637) DE EP 193570 R: AT BE CH DE FR GB IT LI LU NL SE JP 62500289 W 19870205 (198711) C 19880511 (198819) B 19891129 (198948) DE 3518871 EP 193570 R: AT BE CHIDE FR GB IT LI LU NL SE Ġ 19900104 (199003) DE 3574455 US 5089122 (A) 19920218 (199210) DE 3518871 A DE 1985-3518871 19850524; WO 8601425 A WO 1985-DE304 19850902; EP 193570 A EP 1985-904424 19850902; JP 62500289 W JP 1985-504004 19850902; US 5089122 A US 1991-649223 19910128 PRAI DE 1984-257540 19840831; DE 1985-3518871 19850524 3518871 A UPAB: 19930922 A multi-component fluid mixt. flows past one face of one or more membranes at whose other face is maintained (e.g. by suction), a lower partial pressure of one of the components than in the mixt., so causing that component to migrate through the membrane, which may be porous (ultrafiltration) or of pervaporation type. The membrane is heated electrically. It may itself be conductive, or carry a conductive coating, or be inductively heated. Alternatively the membrane is backed by a porous metal support, possibly reinforced by metal netting. The face adjacent the mixt. may be coated with hydrophobic or hydrophilic material, depending on properties of target substance. USE/ADVANTAGE - Suitable e.g. for aq. soln.; oil/water mixt.:bio-reaction prod. or for gaseous mixt. Prevents fluid being cooled by loss of evapn. heat, but highly localised effect does not mar main fluid body. 0/3 L14 ANSWER 20 OF 20 WPIDS COPYRIGHT 2002 DERWENT INFORMATION LTD 1968-06690Q [00] ΑN WPIDS ΤI Desalination wing vinyl pyrrolidone copolymer ion. DC (PURQ) PURAQ CO 1 PΑ CYC

US 3386912 PΙ

(196800) \*

PRAI US 1965-423976 19650107

AB 3386912 A UPAB: 19930831

Low energy desalination of sea water by heating to 48 deg.C in that exchanger and passing it to a chamber where it contacts ion-exchange membrane which has a partly water-sol. particulate resin flowing at 50-52 deg.C through chamber. Water is adsorbed by the resin and separated from the polymer soln. in settling tank.

Copolymer of a) hydrophilic and b) mechanically strengthening hydrophobic monomers. a) is vinyl pyrrolidone and b) is vinyl acetate, MMA, chloroprene, acrylonitrile or pref. a mixt. of styrene and butadiene (partic. 60% vinyl pyrrolidone, 20% styrene, 19.5% butadiene and 0.5% cross-linking trialkyl cyanurate moulded between 'Teflon'-coated glass plates 1/32 in apart). Alternatively a p.v. pyrrolidone membrane coated with an

ion-exchange resin may be used.

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=> fil biosid
'BIOSAD' IS NOT A VACID FILE NAME
SESSION CONTINUES IN FILE 'BIOSIS'
=> fil biosis
FILE 'BIOSIS' ENTERED AT 10:21:13 ON 07 MAR 2002
COPYRIGHT (C) 2002 BFOLOGICAL ABSTRACTS INC. (R)
FILE COVERS 1969 TOIDATE.
CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT
FROM JANUARY 1969 TO DATE.
RECORDS LAST ADDED 6 March 2002 (20020306/ED)
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      (FILE 'WPIDS' ENTERED AT 10:07:39 ON 07 MAR 2002)
                 DEL HIS Y
     FILE 'BIOSIS' ENTERED AT 10:13:47 ON 07 MAR 2002
L1
          772163 S MEMBRANE#
L2
           13439 S (E1) (4A) (DISRUPT? OR ALTER? OR ENHANC? (3A) (TRANSP? OR PERME
L3
          236030 S POLYMER?
L4
             262 S L2 AND L3
L5
          977182 S TRANSPORT?
L6
              45 S 14 AND L5
          482259 S MEMBRANE#/IT
L7
rs
              25 S L6 AND L7
L9
              70 S DISRUPTIVE (3A) AGENT#
L10
               1 S L6 AND L9
L11
               2 S L1 AND L3 AND L9
L12
           20436 S HYDROPHIL?
L13
           48724 S HYDROPHOB?
L14
               4 S L4 AND L12 AND L13
L15
              29 S 18 OR L10 OR L11 OR L14
     FILE 'BIOSIS' ENTERED AT 10:21:13 ON 07 MAR 2002
=> d bib ab it 1-29
L15
     ANSWER 1 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     2002:161277 BIOSIS
AN
DN
     PREV200200161277
TΤ
     PYK2 as a mediator of endothelin-1/Galphall signaling to GLUT4 glucose
     transporters.
     Park, Jin G.; Bose, Avirup; Leszyk, John; Czech, Michael P. (1) (1) Program in Molecular Medicine, 373 Plantation St., Worcester, MA,
AU
     01605: Michael Czech@umassmed.edu USA
SO
     Journal of Biological Chemistry, (December 21, 2001) Vol. 276, No. 51, pp.
     47751-47754. http://www.jbc.org/. print.
     ISSN: 0021-9258.
DT
     Article
LA
     English
     Endothelin-1 (#1-1) signaling through Galphaq/11 stimulates translocation
     of intracellular GLUT4 glucose transporters to the plasma
     membrane of 3\hat{r}\hat{s}+1 adipocytes by an unknown mechanism that requires
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protein tyrosine phosphorylation and ADP-ribosylation factor 6 (ARF6) but

is independent of phosphatidylinositol 3 (PI3)-kinase. In contrast,

insulin action on this process requires PI3-kinase but not ARF6. Here we report the identification of two proteins selectively tyrosinephosphorylated in response to ET-1 but not insulin: the Ca2+-activated tyrosine kinase PYK2 and its physiological substrate, the adhesion scaffold protein paxillin. Endogenous paxillin as well as expressed Myc-tagged PYK2 or a Myc-tagged kinase-deficient PYK2 protein were acutely directed to factin-rich adhesion sites from the adipocyte cytoplasm in response to  $\vec{E}[\vec{T},\vec{T}]$  but not insulin. CADTK-related non-kinase (CRNK) is a dominant negative form of PYK2 containing the C-terminal portion of the protein, which binds paxillin but lacks the PYK2 autophosphorylation site (Tyr402). CRNK expression in 3T3-L1 adipocytes inhibited ET-1-mediated F-actin polymerization and translocation of Myc-tagged GLUT4-enhanced green fluorescent protein (EGFP) to the plasma membrane without disrupting insulin action on these processes. These data reveal the tyrosine kinase PYK2 as a required signaling element in the regulation of GLUT4 recycling in 3T3-L1 adipocytes by ET-1, whereas insulin signaling is directed through a different pathway. Major Concepts Biochemistry and Molecular Biophysics; Cell Biology; Endocrine System (Chemical Coordination and Homeostasis) Parts, Structures, & Systems of Organisms plasma membranes Chemicals & Biochemicals ADP-ribosylation factor 6: analysis, functions, signaling; F-actin: analysis, functions, signaling; GLUT4 glucose transporters: analysis, functions; PYK2: analysis, functions; endothelin-1/G-alpha-11: analysis, functions, signaling; enzymes: analysis, functions; green fluorescentifrotein; insulin: biological activities, functions; kinases: analysis, functions; membrane proteins: analysis, functions paxillin; proteins: analysis, functions; tyrosine: phosphorylation Miscellaneous Descriptors insulin signalling pathways/mechanisms: analysis, functions ORGN Super Taxa Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name 3T3-L1 cell line (Muridae) ORGN Organism Superterms Animals; Chordates; Mammals; Nonhuman Mammals; Nonhuman Vertebrates; Rodents; Vertebrates 9004-10-8 (INSULIN) 9031-44-1 (KINASES) 57186-25-1 (PAXILLIN) 60-18-4Q (TYROSINE) 556-03-6Q (TŸŘÔSINE) ANSWER 2 OF 20 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 2002:23609 BIOSIS PREV200200023609 Method and composition for enhancing transport across biological membranes. Rothbard, Jonathan B. (1); Wender, Paul A. (1) Woodside, CA USA ASSIGNEE: The Board of Trustees of the Leland Stanford, Jr. University, Stanford, CA, USA US 6306993 October 23, 2001 Official Gazette of the United States Patent and Trademark Office Patents, (Oct. 23, 2001) Vol. 1251, No. 4, pp. No Pagination. e-file. ISSN: 0098-1133.

IT

IT

IT

IT

RN

L15 AN

DN TΙ

ΑU

CS

PΤ

SO

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DT
     Patent
LA
     English
AB
     Methods and compositions for transporting drugs and
     macromolecules across biological membranes are disclosed. In one
     embodiment, the invention includes a method for enhancing
     transport of a selected compound across a biological membrane,
     wherein a biological membrane is contacted with a conjugate containing a
     biologically active agent that is covalently attached to a
     transport polymer. In one embodiment, the
     polymer consists of from 6 to 25 subunits, at least 50% of which
     contain a quanidino or amidino sidechain moiety. The polymer is
     effective to impart to the attached agent a rate of trans-membrane
     transport across a biological membrane that is greater than the
     rate of trans-membrane transport of the agent in nonconjugated
ΙT
     Major Concepts
        Biochemistry and Molecular Biophysics; Membranes (Cell
        Biology)
IT
     Chemicals & Biochemicals
        compositions; drugs: pharmaceutical, transport;
        macromolecules: transport
     Miscellaneous Descriptors
IT
        biological membranes
     ANSWER 3 OF 29; BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
AN'
     2002:8763 BIOSIS
DN
     PREV200200008763
     Application of surface modified polypropylene membranes to an anaerobic
TI
     membrane bioreactor.
     Sainbayar, A. (1); Kim, J. S. (1); Jung, W. J. (1); Lee, Y. S. (1); Lee,
ΑU
     (1) School of Chemical Engineering, Seoul National University, Gwanak-qu,
CS
     Sillim-dong, Seoul, 151-744 South Korea
     Environmental Technology, (September, 2001) Vol. 22, No. 9, pp. 1035-1042.
SO
     print.
     ISSN: 0959-3330.
DT
     Article
LA
     English
AB ·
    In order to increase hydrophilicity and thereby to reduce
     membrane fouling caused by hydrophobic adsorption, the surface
     of a hydrophobic 0.2 mum polypropylene (PP) membrane was
     modified by ozone treatment followed by graft polymerization
     with 2-hydroxy-ethyl methacrylate (HEMA). The modified PP (MPP) membranes
     were characterized in terms of contact angle, morphology and degree of
     grafting (DG) The contact angle was reduced from 112degree for a PP
     membrane to nearly Odegree for MPP membranes by introducing functional
     groups such as hydroxyl (-OH) and carbonyl groups (C=O) on the membrane
     surface. As the DG increased, the O/C ratio and membrane resistance of the
     MPP membrane indreased. Using the MPP membrane in the crossflow operation
     of an anaerobic membrane bioreactor (MBR), the membrane
    permeability was enhanced although it was largely dependent on the DG of MPP.
IT
    Major Concepts
        Biomaterials, Bioprocess Engineering, Methods and Techniques
IT
     Chemicals & Biochemicals
        2-hydroxy-ethyl methacrylate [HEMA]; carbonyl group; hydroxyl group
TΤ
    Methods & Equipment
        anaerobic membrane bioreactor: laboratory equipment; crossflow
        operation: production method; graft polymerization:
        production method; ozone treatment: production method
```

Tran 09/755,701 Miscellaneous ΙT hydrophilicity; hydrophobic absorption: membrane fouling effect; membrane permeability; polypropylene membrane: application factorization, contact angle, grafting degree, morphology surface modified RN 868-77-9 (2-HYDROXY-ETHYL METHACRYLATE) ANSWER 4 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15 AN 2001:543445 BIOSIS DN PREV200100543445 ΤI Phospholipid alterations in hepatocyte membranes and transporter protein changes in cholestatic rat model. ΑU Hyogo, Hideyuki; Tazuma, Susumu (1); Nishioka, Tomoji; Ochi, Hidenori; Yamaguchi, Atushi; Numata, Yoshihiro; Kanno, Keishi; Sakomoto, Minoru; Asamoto, Yasumasa; Tsuboi, Kazuhiko; Nakai, Kuniharu; Yasumiba, Shigeyuki; Sunami, Yasushi; Kajiyama, Goro CS (1) First Department of Internal Medicine, Hiroshima University School of Medicine, 1-2#3 Kasumi, Minami-ku, Hiroshima, 734-8551 Japan Digestive Diseases and Sciences, (October, 2001) Vol. 46, No. 10, pp. SO 2089-2097. print. ISSN: 0163-2116 DTArticle LA English SLEnglish AΒ Biliary components are transported by hepatic adenosine triphosphate binding cassette (ABC) transporters that are located in canalicular membranes. Physiological transporter function is related to membrane fluidity, which is modulated by the phospholipid cómposition of the lipid bilayer. We hypothesized that cholestasis may alter transporter function by modifying phospholipid species to protect the cell from cholestatic damage.

Therefore, we sexamined the expression of ABC transport proteins and their mRNA levels in canalicular membrane vesicles isolated from rat liver 6 hr or three days after bile duct ligation. Membrane lipid composition and membrane fluidity of both sinusoidal and canalicular membrane vesicles were also examined. By 6 hr after bile duct ligation, we found a clear increase of mdr2 and bsep mRNA. These changes were associated with an increase of mdr-Pgp and with a clear decrease of mrp2 protein, and small decrease of bsep protein. In addition, mdrlb mRNA showed a strong increase by three days after bile duct ligation. Canalicular membrane fluidity decreased in a marked time-dependent manner, whereas sinusoidal membranes showed biphasic changes: increased fluidity at 6 hr and a decrease at three days. These changes were closely related to the changes of membrane lipid constitution; the saturated/unsaturated fatty acid ratio increased for phosphatidylcholine in canalicular membrane and the reverse occurred in sinusoidal membrane, and those for sphingomyelin showed the opposite pattern. We conclude that cholestasis causes modulation of ABC transporters as well as that of the lipid constituțion in lipid bilayer. These may confer cytoprotective resistance to hepatocytes against cholestatic stress.

IT Major Concepts

Biochemistry and Molecular Biophysics; Digestive System (Ingestion and Assimilation); Membranes (Cell Biology)

IT Parts, Structures, & Systems of Organisms

bile duct: digestive system; hepatocytes: digestive system, membranes; liver: digestive system

IT Diseases

cholestasis digestive system disease, histopathology

IT Chemicals & Biochemicals

ATP-binding cassette transporters: expression; besp messenger

```
RNA; bsep protein; lipid bilayer; mdrlb messenger RNA; mdr2 messenger
        RNA; membrane phospholipids: alterations; mrp2
        protein; phosphatidylcholine; sphingomyelin; transporter
        proteins
     Alternate Indexing
IT
        Cholestasis (MeSH)
IΤ
     Methods & Equipment
        Western blot: detection method, gene mapping, labeling; bile duct
        ligation: surgical method; fluorescence polarization analysis:
        analytical method; reverse transcriptase-polymerase chain
        reaction: genetic method, polymerase chain reaction
IT
     Miscellaneous Descriptors
          membrane fluidity
ORGN Super Taxa
        Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        rat (Muridae): animal model, strain-Sprague-Dawley
ORGN Organism Superterms
        Animals; Chordates; Mammals; Nonhuman Mammals; Nonhuman Vertebrates;
        Rodents; Vertebrates
    ANSWER 5 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
ΑN
     2001:48510 BIOSIS
     PREV200100048510
DN
     pH-responsive pseudo-peptides for cell membrane
TΙ
     disruption.
     Eccleston, M. E.; Kuiper, M.; Gilchrist, F. M.; Slater, N. K. H. (1)
ΑU
CS
     (1) Departmen of Chemical Engineering, University of Cambridge, Pembroke
     Street, Cambridge, CB2 3RA: nigel slater@cheng.cam.ac.uk UK
SO
     Journal of Controlled Release, (3 November, 2000) Vol. 69, No. 2, pp.
     297-307. print!
     ISSN: 0168-3659.
DT
     Article
LA
     English
SL
     English
     We describe pseudo-peptides obtained by the copolymerisation of L-lysine
AΒ
     and L-lysine ethyl-ester with various hydrophobic dicarboxylic
     acid moieties. In aqueous solution, when the carboxylic acid groups are
     charged, the polymers dissolve. When they are fully neutralised
     the hydrophobic moieties cause the polymer to
     precipitate. The pH range over which reversible precipitation occurs can
     be adjusted by changing the intramolecular hydrophilic/
    hydrophobic balance, by using a carboxylic acid moiety with a
     different pka value or by changing the apparent pka value of the
    polymer through chemical modifications of the backbone. These
     bio-degradable materials are well tolerated by a range of mammalian cell
     lines at physiclogical pH but display an ability to associate with the
     outer membranes of these cells, which they rupture to varying degrees at
     pH 5.5. Relatiÿ́e to the degree of lysis displayed by poly(L-lysine
     iso-phthalamide), lysis was reduced by partial esterification and
     increased by replacing the aromatic iso-phthaloyl moiety with a long chain
     aliphatic dodecyl moiety. Similar behaviour was observed for the
    pH-dependent rupture of human erythrocytes, where poly(L-lysine
     dodecanamide) displayed enhanced cell lysis at pH values <7.0 relative to
    poly(L-lysine iso-phthalamide).
    Major Concepts
ΙT
        Biochemistry and Molecular Biophysics; Membranes (Cell Biology)
ΙT
     Parts, Structures, & Systems of Organisms
        erythrocytes: blood and lymphatics
     Chemicals & Biochemicals
ΙT
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L-lysine; Lilysine ethyl ester; poly(L-lysine dodecanamide)
IT
     Miscellaneous Descriptors
        cell membrane disruption
ORGN Super Taxa
        Hominidae primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae)
ORGN Organism Superterms
        Animals; Chordates; Humans; Mammals; Primates; Vertebrates
     56-87-1 (L-LYSINE)
RN
     4117-33-3 (L≟ĒŸSINE ETHYL ESTER)
    ANSWER 6 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
     2000:533813 BIOSIS
ΑN
     PREV200000533813
DN
TI
     Membranes for endotoxin removal from dialysate: Considerations on
     feasibility of commercial ceramic membranes.
ΑU
     Bender, Heiko Eflaenzel, Anne; Saunders, Nicola; Czermak, Peter (1);
     Catapano, Gerardo; Vienken, Joerg
     (1) Department KMUB-Biotechnology, University of Applied Sciences,
CS
     Institute of Biochemical Engineering and Membrane Technology,
     Wiesenstrasse 14, D-35390, Giessen Germany
     Artificial Organs, (October, 2000) Vol. 24, No. 10, pp. 826-829. print.
SO
     ISSN: 0160-564X.
DT
     Article
     English
LA
SL
     English
AB
     As the quality of water in dialysis fluid varies considerably, dialysate
     is often contaminated by large amounts of bacteria and endotoxins.
     Membrane properties and operating pressures are acknowledged to give
     high-flux dialivsis with bicarbonate the bacteriological potential to favor
    passage of endotoxin fragments from the dialysate into the blood stream.
     Therefore, a sterile dialysate will have to become a standard.
     Ultrafiltration across hydrophobic synthetic membranes was shown to remove
     endotoxins (and their fragments) from dialysis water by the combined
     effect of filtration and adsorption. However, each module can be used for
     a limited time only. Ceramic membranes may represent an
     alternative to polymeric membranes for
     endotoxin removal. In this article, we tested the capacity of different
     commercial ceramic membranes with nominal molecular weight cut-off down to
     1,000 to retain endotoxins from Ps. aeruginosa. The tested membranes did
    not generally produce dialysate meeting the Association for the
     Advancement of Medical Instrumentation standard. When using
     aluminum-containing membranes, we detected aluminum leaking into the
     dialysate that could possibly be transported into the blood
    stream.
ΙT
    Major Concepts
        Biomaterials
IT
    Chemicals & Biochemicals
        Pseudomonas aeuroginosa endotoxin; endotoxin: dialysate removal
TΤ
    Miscellaneous Descriptors
       blood stream; commercial ceramic membrane; dialysis fluid:
       water quality; hydrophobic synthetic membrane;
       membrane: endotoxin removal, operating pressure
ORGN Super Taxa
        Pseudomonadaceae: Gram-Negative Aerobic Rods and Cocci, Eubacteria,
       Bacteria, Microorganisms
ORGN Organism Name
        Pseudomonas (Pseudomonadaceae)
ORGN Organism Superterms
```

#### Bacteria; Eubacteria; Microorganisms

```
L15
     ANSWER 7 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     2000:504696 BÍOSIS
AN
     PREV200000504696
DN
     De novo synthesis of proteinase 3 by cytokine primed circulating human
ΤI
     polymorphonuclear neutrophils and mononuclear cells.
     Zhou, Zhijie; Richard, Carol; Menard, Henri A. (1)
ΑU
     (1) Division of Rheumatology, McGill University Health Centre, 1650 Cedar Ave., A6-162.1; Montreal, PQ, H3G 1A4 Canada
CS
     Journal of Rhedmatology, (October, 2000) Vol. 27, No. 10, pp. 2406-2411.
SO
     print.
     ISSN: 0315-162X.
DΤ
     Article
                 1 4
     English
LA
SL
     English
AB
     Objective: When polymorphonuclear neutrophils (PMN) and peripheral blood
     monocytes (PBMC) are stimulated with tumor necrosis factor alpha
     (TNF-alpha), preexisting granule stored proteinase 3 (PR3) is translocated
     to the surface of their plasma membrane. We investigated whether PR3 gene
     reactivation and new PR3 protein production were also features of priming
     by cytokine. Methods: Normal human PMN and PBMC were isolated and
     stimulated in the tro with TNF-alpha. They were harvested at different
     intervals and subjected to total RNA and protein analysis. PR3 mRNA was
     identified by reverse transcription polymerase chain reaction,
     Northern blot, and sequencing. De novo PR3 synthesis was evaluated by
     metabolic labeling with (35S) methionine followed by immunoprecipitation
     using anti-neutrophil cytoplasmic antibodies from serum of patients with
     active Wegener's granulomatosis and mouse monoclonal anti-native PR3 antibodies. Results: Resting PMN and PBMC do not express PR3 mRNA. During
     priming, PR3 MRNA appears in PMN at 2 h, peaks at 6 h, and has disappeared
     at 12 h. By comparison, in primed PBMC, PR3 mRNA appears at 6 h, peaks at
     12 h, and disappears at 24 h. Immunoprecipitation of metabolically labeled
     PR3 revealed new synthesis of PR3 by both cell types, a process that was
     inhibited by cycloheximide. Conclusion: Primed PMN and PBMC can express
     PR3 mRNA and synthesize new PR3 protein, providing an alternative
     source to membrane PR3. Whether that small amount of inducible
     PR3 has a primăry structure, a localization, or a role different from
     those of prefermed PR3 stored in granules remains to be clarified.
IT
     Major Concepts
        Cell Biology; Immune System (Chemical Coordination and Homeostasis);
        Blood and Tymphatics (Transport and Circulation)
IT
     Parts, Structures, & Systems of Organisms
        peripheral blood monocytes: blood and lymphatics, circulating, cytokine
        primed, immune system; plasma membrane; polymorphonuclear
        neutrophilisi blood and lymphatics, circulating, cytokine primed, immune
        system
     Chemicals & Biochemicals
IΤ
        cyclohexamide; proteinase 3: de novo synthesis; tumor necrosis factor
        alpha
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae)
ORGN Organism Superterms
        Animals; Chordates; Humans; Mammals; Primates; Vertebrates
RN
     128028-50-2 (PROTEINASE 3)
```

ANSWER 8 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

Page 61

2000:329357 BIOSIS

L15

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PREV200000329357
DN
     Hgtlp, a high affinity glutathione transporter from the yeast Saccharomyces gerevisiae.
ΤI
     Bourbouloux, Andree; Shahi, Puja; Chakladar, Abhijit; Delrot, Serge;
ΑU
     Bachhawat, Anang K. (1)
(1) Institute of Microbial Technology, Sector 39-A, Chandigarh, 160 036
CS
     India
SO
     Journal of Biological Chemistry, (May 5, 2000) Vol. 275, No. 18, pp.
     13259-13265. print.
     ISSN: 0021-9258.
DT
     Article
     English
LA
SL
     English
     A high affinity glutathione transporter has been identified,
AΒ
     cloned, and characterized from the yeast Saccharomyces cerevisiae. This
     transporter, Hgtlp, represents the first high affinity glutathione
     transporter to be described from any system so far. The strategy
     for the identification involved investigating candidate glutathione
     transporters from the yeast genome sequence project followed by
     genetic and physiological investigations. This approach revealed HGT1
     (open reading frame YJL212c) as encoding a high affinity glutathione transporter. Yeast strains deleted in HGT1 did not show any
     detectable plasma membrane glutathione transport, and
     hgt1DELTA disruptants were non-viable in a glutathione
     biosynthetic mutant (gsh1DELTA) background. The glutathione repressible
     transport activity observed in wild type cells was also absent in
     the hgt1DELTA strains. The transporter was cloned and kinetic
     studies indicated that Hgtlp had a high affinity for glutathione (Km = 54
     muM)) and was hot sensitive to competition by amino acids, dipeptides, or
     other tripeptiges. Significant inhibition was observed, however, with
     oxidized glutathione and glutathione conjugates. The transporter reveals a novel class of transporters that has homologues in other yeasts and plants but with no apparent homologues in either
     Escherichia coll or in higher eukaryotes other than plants.
ΙT
     Major Concepts
         Molecular Genetics (Biochemistry and Molecular Biophysics); Methods and
         Techniques?
IT
     Parts, Structures, & Systems of Organisms
         plasma membrane
ΙT
     Chemicals & Biochemicals
         Hgtlp: high affinity glutathione transporter; glutathione
         transporters; Saccharomyces cerevisiae HGT1 gene (Ascomycetes)
ΙT
     Methods & Equipment
        DNA isolation: Extraction, Isolation, Purification and Separation Techniques, isolation method; PCR [polymerase chain reaction] DNA amplification, DNA amplification method, in-situ
         recombinantigene expression detection, sequencing techniques; cloning:
         Recombinan PNA Technology, cloning method; kinetic analysis: activity
         assays, analytical method; synthesis: Synthetic Techniques, synthetic
         method; tetrad analysis: Molecular Biology Techniques and Chemical
         Characterization, analytical method
ΙT
     Miscellaneous Descriptors
         amino acid sequence
ORGN Super Taxa
         Ascomycetes Fungi, Plantae
ORGN Organism Name
         Saccharomyces cerevisiae (Ascomycetes)
ORGN Organism Superterms
```

Fungi; Microorganisms; Nonvascular Plants; Plants

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ANSWER 9 OF 29 BIOSIS
L15
                             COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     2000:279411 BIOSIS
AN
DN
     PREV200000279411
     Expression of aquaporin-4 water channels in rat cholangiocytes.
ΤI
     Marinelli, Raul A.; Pham, Linh D.; Tietz, Pamela S.; LaRusso, Nicholas F.
ΑU
     (1)
CS
     (1) Center for Basic Research in Digestive Diseases, Mayo Clinic, 200
     First Street SW, Rochester, MN, 55905 USA
     Hepatology, (June, 2000) Vol. 31, No. 6, pp. 1313-1317. print.
SO
     ISSN: 0270-9139.
DT
     Article
     English
LA
ŞL
     English
     We recently reported that secretin induces the exocytic insertion of
AB
     functional aquaporin-1 water channels (AQP1) into the apical membrane of
     cholangiocytes and proposed that this was a key process in ductal bile
     secretion. Because AQP1 is present on the basolateral cholangiocyte
     membrane in low amounts, we hypothesized that another AQP must be
     expressed at this domain to facilitate transbasolateral water movement.
     Thus, we investigated the expression, subcellular localization, possible
     regulation by secretin, and functional activity of AQP4, a
     mercury-insensitive water channel expressed in other fluid
     transporting epithelia. Using reverse transcription-
     polymerase chain reaction (RT-PCR) on RNA prepared from purified
     rat cholangiocytes, we amplified a product of 311 bp that was 100%
     homologous to the reported AQP4 sequence. RNase protection assay confirmed
     the presence of an appropriate size transcript for AQP4 in cholangiocytes.
     Immunoblotting detected a band of approximately 31 kd corresponding to
     AOP4 in basolateral but not apical membranes of cholangiocytes. Secretin
     did not alter the amount of plasma membrane AQP4 but,
     as expected, induced AQP1 redistribution from intracellular to apical
     plasma membranes. Functional studies showed that AQP4 accounts for about
     15% of total cholangiocyte membrane water permeability. Our results
     indicate that: (1) cholangiocytes express AQP4 messenger RNA (mRNA) and protein and (2) in contrast to AQP1, which is targeted to the apical
     cholangiocyte membrane by secretin, AQP4 is constitutively expressed on
     the basolateral cholangiocyte membrane and is secretin unresponsive. The
     data suggest that AQP4 facilitates the basolateral transport of
     water in cholangiocytes, a process that could be relevant to ductal bile
     formation.
     Major Concepts:
TΤ
        Molecular Genetics (Biochemistry and Molecular Biophysics);
        Membranes (Cell Biology); Digestive System (Ingestion and
        Assimilation)
IT
     Parts, Structures, & Systems of Organisms
        cholangiocyte
TT
     Chemicals & Biochemicals
        RNA; RNase protection assay; aquaporin-4 water channel messenger RNA;
        aquaporin-4 water channels; secretin
ΙT
     Methods & Equipment
        immunoblotting; reverse transcriptase-polymerase chain
        reaction
ΙT
     Miscellaneous Descriptors
        transmembrane water transport
ORGN Super Taxa
        Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name (Muridae)
ORGN Organism Superterms
        Animals; Chordates; Mammals; Nonhuman Mammals; Nonhuman Vertebrates;
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Rodents; Vertebrates
RN
    1393-25-5 (SECRETIN)
    ANSWER 10 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
    2000:268789 BIOSIS
AN
DN
    PREV200000268789
    Phospholipid alterations in rat hepatocyte membranes
ΤI
    in cholestasis and cytoprotective transporter protein changes.
    Hyogo, Hideyuki (1); Tazuma, Susumu; Sakamoto, Minoru; Nakai, Kuniharu;
ΑU
    Asamoto, Yasumasa; Tsuboi, Kazuhiko; Yasumiba, Shigeyuki; Sunami, Yasushi;
    Ochi, Hidenori Muller, Michael; Kajiyama, Goro
CS
     (1) Hiroshima Univ, Hiroshima Japan
    Gastroenterology, (April, 2000) Vol. 118, No. 4 Suppl. 2 Part 1, pp. AASLD
SO
    Association and the Digestive Disease Week. San Diego, California, USA May
    21-24, 2000 American Gastroenterological Association
     . ISSN: 0016-5085.
    Conference
DT
    English
LΑ
    English
SL
IT
    Major Concepts
         Membranes (Cell Biology); Digestive System (Ingestion and
       Assimilation)
    Parts, Structures, & Systems of Organisms
ΙT
       hepatocyte membranes
IT
    Diseases
       cholestasis digestive system disease
IT
    Chemicals & Bijochemicals
       adenosine-triphosphate-binding cassette; cytoprotective
       transporter protein; phospholipids: alterations
IT
    Alternate Indexing
       Cholestasis (MeSH)
    Methods & Equipment
ΙT
       RT-PCR [reverse transcriptase-polymerase chain reaction]:
       amplification method, analytical method; Western blotting: analytical
       method
ΙT
    Miscellaneous Descriptors
       Meeting Abstract
ORGN Super Taxa
       Muridae: Rödentia, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
       rat (Muridae)
ORGN Organism Superterms
       Animals; Chordates; Mammals; Nonhuman Mammals; Nonhuman Vertebrates;
       Rodents; Vertebrates
L15 ANSWER 11 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
AN
    2000:79036 BIOSIS
DN
    PREV200000079036
TI
    Functional characterization of mutations in melanocortin-4 receptor
    associated with human obesity.
ΑU
    Ho, Guyu; MacKenzie, Robert G. (1)
    (1) Dept. of Cell Biology, Parke-Davis Pharmaceutical Research, 2800
CS
    Plymouth Rd., Ann Arbor, MI USA
SO
    Journal of Biological Chemistry, (Dec. 10, 1999) Vol. 274, No. 50, pp.
    35816-35822.
    ISSN: 0021-9258
DT
    Article
LA
    English
```

SL Melanocortin-4 receptor (MC4R) is a G protein-coupled receptor implicated AΒ in the regulation of body weight. Genetic studies in humans have identified two frameshift mutations of MC4R associated with a dominantly inherited form of obesity. We have generated and expressed the corresponding MC4R mutants in 293T cells and found that cells transfected with the truncation mutants failed to exhibit agonist binding or responsiveness despite retention of structural motifs potentially sufficient for binding and signaling. Immunofluorescence studies showed that the mutant proteins were expressed and localized in the intracellular compartment but absent from the plasma membrane, suggesting that these mutations disrupted the proper cellular transport of MC4R. Further studies identified a sequence in the cytoplasmic tail of MC4R necessary for the cell surface targeting. We further investigated a possible dominant-negative activity of the mutants on wild-type receptor function. Co-transfection studies showed that the mutants affected neither signaling nor cell surface expression of wild-type MC4R. We also characterized three human sequence variants of MC4R, but these exhibited identical affinities for peptide ligands and identical agonist responsiveness! Thus, unlike the obesity-associated MC4R truncation mutants, the polymorphisms of MC4R are unlikely to be contributors to human obesity Major Concepts IT Molecular Genetics (Biochemistry and Molecular Biophysics); Methods and Techniques Parts, Structures, & Systems of Organisms IT plasma membrane IT Diseases obesity: nutritional disease Chemicals & Biochemicals IT G protein; melanocortin-4 receptor Alternate Indexing IT Obesity (MeSH) IT Methods & Equipment PCR [polymerase chain reaction]: DNA amplification, DNA amplification method, in-situ recombinant gene expression detection, sequencing techniques; adenylate cyclase assay: Analysis/Characterization Techniques: CB, analytical method; cAMP iodine-125 scintillation proximity assay system: Amersham Pharmacia Biotech, laboratory equipment; cell culture: Cell Culture Techniques, cell culture method; cloning: Recombinant DNA Technology, cloning method; confocal laser scanning microscope: Olympus, laboratory equipment; immunofluorescence: microscopy method, microscopy: CB; oligonucleotide-directed mutagenesis: genetic method, mutagenesis, protein engineering; receptor binding assay: analytical method, binding assays; transfection: gene expression/vector techniques, genetic method IT Miscellaneous Descriptors body weight; frameshift mutation; gene polymorphism ORGN Super Taxa Cercopithe didae: Primates, Mammalia, Vertebrata, Chordata, Animalia; Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name 293T cell line (Hominidae); COS-7 cell line (Cercopithecidae); human (Hominidae) ORGN Organism Superterms

L15 ANSWER 12 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. AN 2000:49083 BIOSIS

Primates; Nonhuman Vertebrates; Primates; Vertebrates

Animals; Chordates; Humans; Mammals; Nonhuman Mammals; Nonhuman

```
Tran 09/755,701
DN
     PREV200000049083
     Cytosolic delivery of granzyme B by bacterial toxins: Evidence that
ΤI
     endosomal distuption, in addition to transmembrane pore formation, is an
     important function of perforin.
     Browne, Kylie A; Blink, Elizabeth; Sutton, Vivien R.; Froelich, Christopher J; Jans, David A.; Trapani, Joseph A. (1)
(1) Austin Research Institute, Studley Road, Heidelberg, VIC Australia
ΑU
CS
     Molecular and cellular Biology, (Dec., 1999) Vol. 19, No. 12, pp.
SO
     8604-8615.
     ISSN: 0270-7306.
DT
     Article
LA
     English
SL
     English
     Granule-mediated cell killing by cytotoxic lymphocytes requires the
AΒ
     combined actions of a membranolytic protein, perforin, and
     granule-associated granzymes, but the mechanism by which they jointly kill
     cells is poorly understood. We have tested a series of membrane-
     disruptive agents including bacterial pore-forming
     toxins and hemolytic complement for their ability to replace perforin in
     facilitating granzyme B-mediated cell death. As with perforin, low
     concentrations of streptolysin O and pneumolysin (causing <10% 51Cr
     release) permitted granzyme B-dependent apoptosis of Jurkat and Yac-1
     cells, but staphylococcal alpha-toxin and complement were ineffective,
     regardless of concentration. The ensuing nuclear apoptotic damage was
     caspase dependent and included cleavage of poly(ADP-ribose)
     polymerase, suggesting a mode of action similar to that of
     perforin. The plasma membrane lesions formed at low dose by
     perforin, pneumolysin, and streptolysin did not permit diffusion of
     fluorescein-labeled proteins as small as 8 kDa into the cell, indicating
     that large membrane defects are not necessary for granzymes (32)
     to 65 kDa) to enter the cytosol and induce apoptosis. The endosomolytic
     toxin, listeriolysin O, also effected granzyme B-mediated cell death at
     concentrations which produced no appreciable cell membrane
     damage. Cells pretreated with inhibitors of endosomal trafficking such as
     brefeldin A took up granzyme B normally but demonstrated seriously
     impaired nuclear targeting of granzyme B when perforin was also added,
     indicating that an important role of perforin is to disrupt vesicular
     protein trafficking. Surprisingly, cells exposed to granzyme B with
     perforin concentrations that produced nearly maximal 51Cr release (1,600
     U/ml) also underwent apoptosis despite excluding a 8-kDa
     fluorescein-labeled protein marker. Only at concentrations of >4,000 U/ml
     were perforin pores demonstrably large enough to account for transmembrane
     diffusion of granzyme B. We conclude that pore formation may allow
     granzyme B direct cytosolic access only when perforin is delivered at very
     high concentrations, while perforin's ability to disrupt endosomal
     trafficking may be crucial when it is present at lower concentrations or
     in killing cells that efficiently repair perforin pores.
ΙT
     Major Concepts
        Enzymology Biochemistry and Molecular Biophysics); Membranes
```

(Cell Biology); Toxicology

Parts, Structures, & Systems of Organisms IT

endosome; plasma membrane: pore

ΙT Chemicals & Biochemicals

granzyme B $\frac{1}{2}$ Listeriolysin: toxin; perforin; pneumolysin; staphylococcal alpha-toxin; streptolysin O

Miscellaneous "Descriptors ΙT

apoptosis

ORGN Super Taxa

Bacteria: Microorganisms; Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia; Muridae: Rodentia, Mammalia, Vertebrata, Chordata,

Animalia ORGN Organism Name Jurkat cell line (Hominidae); Yac-1 cell line (Muridae); bacteria (Bacteria) ORGN Organism Superterms Animals; Bacteria; Chordates; Eubacteria; Humans; Mammals; Microorganisms; Nonhuman Mammals; Nonhuman Vertebrates; Primates; Rodents: Vertebrates 143180-74-9 (GRANZYME B) RNANSWER 13 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15 ΑN 2000:9745 BIOSIS PREV200000009745 DN ΤI The design and synthesis of polymers for eukaryotic membrane disruption. ΑU Murthy, Niren; Robichaud, John R.; Tirrell, David A.; Stayton, Patrick S.; Hoffman, Allan S. (1) CS (1) Department of Bioengineering, University of Washington, Seattle, WA, SO Journal of Controlled Release, (Aug. 27, 1999) Vol. 61, No. 1-2, pp. 137-143. ISSN: 0168-36591 DTArticle -LA English SLEnglish The intracellular trafficking of drugs is critical to the efficacy of AB drugs that are susceptible to attack by lysosomal enzymes. It is therefore an important goal to design and synthesize molecules which can enhance the transport of endocytosed drugs from the endosomal compartments to . the cytoplasm The pH of an endosome is lower than that of the cytosol by one to two phanits, depending on the stage of endosomal development. This pH gradient is a key factor in the design of membranedisruptive polymers which could enhance the endosomal release of drugs. Such polymers should disrupt lipid bilayer membranes at pH 6.5 and below, but should be non-lytic at pH 7.4. We have designed and synthesized pH-sensitive synthetic polymers which efficiently disrupt red blood cells within a sharply defined pH range. One of these polymers, poly(ethyl acrylic acid) (PEAAc) has been previously shown to disrupt synthetic vesicles in a pH-dependent fashion (6). PEAAc hemolyzes red blood cells with an activity of 107 molecules per red blood cell, which is as efficient on a molar basis as the peptide melittin. The mechanism of RBC hemolysis by PEAAc is consistent with the colloid osmotic mechanism. PEAAc's hemolytic activity rises rapidly as the pH decreases from 6.3 to 5.0, and there is no hemolytic activity at pH 7.4. A related polymer, poly propyl acrylic acid) (PPAAc), was synthesized to test whether making the pendant alkyl group more hydrophobic by adding one methylene group would increase the hemolytic activity. PPAAc was found to disrupt red bilod cells 15 times more efficiently than PEAAc at pH 6.1. PPAAc was also not active at pH 7.4 and displayed a pH-dependent hemolysis that was shifted toward higher pH's. Random 1:1 copolymers of ethyl acrylate (EA) and acrylic acid (AAc) (which contain random -COOH and -C2H5 groups that are present and regularly repeat in PEAAc) also displayed significant hemolytic activity, with an efficiency close to PEAAc. These results demonstrate that pH-sensitive synthetic polymers can be molecularly engineered to efficiently disrupt eukaryotic membranes within defined and narrow pH ranges. Thus, these polymers might serve as endosomal disruptive agents with specificities for early or late endosomes.

Major Concepts

TΤ

```
Membranes (Cell Biology); Pharmacology
     Parts, Structures, & Systems of Organisms
ΙT
        endosome; red blood cell: blood and lymphatics; red cell
        membrane: disruption
     Chemicals & Biochemicals
IT
        melittin; poly(propyl acrylic acid); poly(propyl acrylic acid)
IT
     Miscellaneous Descriptors
        drug delivery; pH
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia;
        Hymenoptera Insecta, Arthropoda, Invertebrata, Animalia
ORGN Organism Name
        bee (Hymenoptera); human (Hominidae)
ORGN Organism Superterms
        Animals; Arthropods; Chordates; Humans; Insects; Invertebrates;
        Mammals; Primates; Vertebrates
RN
     20449-79-00 (MEDITTIN)
     37231-28-0Q (MELITTIN)
    ANSWER 14 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
     1999:112240 BIOSIS
ΑN
     PREV199900112240
DN
TΙ
    Alterations in adhesion, transport, and
    membrane characteristics in an adhesion-deficient pseudomonad.
ΑU
     Deflaun, M. F. . Oppenheimer, S. R.; Streger, S.; Condee, C. W.; Fletcher,
    M.(1)
ĊS
     (1) Belle W. Baruch Inst. Marine Biol. Coastal Res., Univ. South Carolina,
     Columbia, SC 29208 USA
SO
    Applied and Environmental Microbiology, (Feb., 1999) Vol. 65, No. 2, pp.
     759-765.
ISSN: 0099-2240
\mathsf{D}\mathbf{T}
    Article
LA
    English
    A stable adhesion-deficient mutant of Burkholderia cepacia G4, a soil
AB
    pseudomonad, was selected in a sand column assay. This mutant (ENV435) was
     compared to the wild-type strain by examining the adhesion of the
     organisms to silica sand and their transport through two aquifer
     sediments that differed in their sand, silt, and clay contents. We
    compared the longitudinal transport of the wild type and the
     adhesion mutant to the transport of a conservative chloride
    tracer in 25 cm long glass columns. The transport of the
    wild-type strain was severely retarded compared to the transport
    of the conservative tracer in a variety of aquifer sediments, while the
    adhesion mutant and the conservative tracer traveled at similar rates. An
    intact sediment core study produced similar results; ENV435 was
     transported at a faster rate and in much greater numbers than G4.
    The results of hydrophobic interaction chromatography revealed that G4 was
    significantly more hydrophobic than ENV435, and polyacrylamide gel
    electrophoresis revealed significant differences in the lipopolysaccharide
    O-antigens of the adhesion mutant and the wild type. Differences in this
    cell surface polymer may explain the decreased adhesion of
    strain ENV435:
ΙT
    Major Concepts
        Cell Biology: Pollution Assessment Control and Management
ΙT
    Parts, Structures, & Systems of Organisms
        cell membranes: characteristics
ΙT
    Chemicals & Biochemicals
        cell surfaçe polymers; O antigens
    Miscellaneous Descriptors
ΙT
        adhesion-defective mutants; aquifer sediments; bacterial adhesion
```

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alterations; transport; transport rates
ORGN Super Taxa
        Pseudomonadaceae: Gram-Negative Aerobic Rods and Cocci, Eubacteria,
        Bacteria, Microorganisms
ORGN Organism Name
        pseudomonadis (Pseudomonadaceae); Burkholderia cepacia
        (Pseudomonadaceae): strain-G4
ORGN Organism Superterms
        Bacteria; Eubacteria; Microorganisms
L15
     ANSWER 15 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
     1998:428583 BLOSIS
ΑN
DN
     PREV199800428583
ΤI
     Active platelet movements on hydrophobic/hydrophilic
     microdomain-structured surfaces.
ΑU
     Ito, Etsuko; Súzuki, Ken; Yamato, Masayuki; Yokoyama, Masayuki; Sakurai,
     Yasuhisa; Okano, Teruo (1)
     (1) Inst. Biomed. Eng., Tokyo Women's Med. Univ., 801 Kawada-cho,
CS
     Shinjuku-ku, Tôkyo 162-8666 Japan
SO
     Journal of Biomedical Materials Research, (Oct., 1998) Vol. 42, No. 1, pp.
     ISSN: 0021-9304.
DT
     Article
LA
     English
     The early motion and interaction of platelets on a microdomain-structured
AΒ
     block copolymer surface composed of 2-hydroxyethyl methacrylate
     (HEMA)-styrene were analyzed and compared with those on a compositionally
     identical random copolymer, homopolymer poly (HEMA) (hydrophilic
     ) and polystyrene (hydrophobic) surfaces. Contacting platelets
     were quantitatively more active, with motions including rolling,
     detachment, oscillatory vibration, and change of direction only on the
     HEMA-St block ීල්ලාolymer surface. Active platelet movements were observed
     for long time periods (>20 min) on HEMA-St block copolymer surfaces and
     were distinct from those for inert PSt latex particles on these same
     surfaces, demonstrating that platelet movements were not due to physical
     forces such as convection, hydrophobic interactions, or
    microbrownian movement. To study the cause and mechanism underlying the
     platelet movements, platelets treated with an adenosine triphosphate (ATP)
     synthesis inhibition, NaN3, or a membrane skeleton-
     disrupting chemical agent, dibucaine, were also studied on these
     surfaces. Both treatments reduced platelet movement and demonstrated that
     platelets in contact with the HEMA-St block copolymer surface require
    metabolic processes consuming ATP and involve dynamics of their membrane
    skeleton. These energy-consuming active movements might explain the
    previously observed lower platelet activation and low thrombogenicity of
     the HEMA-St block copolymers. Enhanced platelet movements on the HEMA-St
    block copolymer surface show that the microdomain surface interacts
     uniquely with platelets to hinder activation and preserve passive platelet
     function despite surface contact.
    Major Concepts:
ΙT
        Biomaterials; Blood and Lymphatics (Transport and
       Circulation)
ΙT
    Chemicals & Brochemicals
         hydrophobic-hydrophilic microdomain-structured
        surfaces; poly(2-hydroxyethyl methacrylate); polystyrene; ATP;
        2-hydroxyethyl methacrylate-styrene copolymer
    Miscellaneous Descriptors
ΙT
       active platelet movements; blood-polymer interaction;
       membrane skeleton dynamics; platelet activation inhibition
```

25249-16-5 (POLY(2-HYDROXYETHYL METHACRYLATE))

117

RN

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9003-53-6 (POLYSTYRENE)
     56-65-5Q (ATP)
     42530-29-0Q (ATP)
     94587-45-8Q (ATP)
     111839-44-20 (ATP)
     ANSWER 16 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
     1998:387929 BIOSIS
AN
DN
     PREV199800387929
ΤI
     Renal Na+, K+-ATPase in SHR: Studies of activity and gene expression.
ΑU
     Nguyen, A.-T: [[[1]]; Hayward-Lester, A.; Sabatini, S.; Doris, P. A.
CS
     (1) Inst. Mola Med., Univ. Tex. Health Sci. Cent., 2121 W. Holcombe Blvd.,
     Houston, TX 77030 USA
SO
     Clinical and Experimental Hypertension, (July-Aug., 1998) Vol. 20, No.
     5-6, pp. 641-656.
     ISSN: 1064-1963.
DT
     Article
LA
     English
     The mechanism by which increased dietary intake of calcium reduces blood
AB
     pressure in the spontaneously hypertensive rat is unknown. The present
     studies were designed to determine if there were alterations in the
     activity of the major membrane ion translocating pump, sodium,
     potassium-ATPase (NKA), in the kidneys of hypertensive rats and whether
     increased dietary calcium intake affected the activity of this enzyme.
     Fifteen-week old SHR's were found to have lower total ATPase activity in
     microsomal preparations from the kidney than age matched Wistar-Kyoto
     animals. Both the ouabain-sensitive component (NKA) and the
     ouabain-insensitive component were lower in SHR. Increasing dietary
     calcium intake from 1% to 3% elevated both components of the ATPase
     activity in SHR, but was without effect in WKY. Measurement of
     membrane phospholipid composition suggested that altered
     phospholipid composition did not account for the reduced ATPase activity
     observed, but indicated a reduced density of ATPase in SHR. A technique
     has been devised for qualitative and quantitative analysis of Na, K-ATPase
     alpha isoforms using RT-PCR. This technique reveals that the alpha I
     isoform is the sole catalytic isoform present in the nephron. Accurate and
     precise quantification of the amount of gene expression in individual
     nephron segments is reported and will be applied to determine whether
     dietary calcium influences blood pressure by a mechanism which alters
     nephron NKA gene expression.
IT
     Major Concepts
        Cardiovascular System (Transport and Circulation); Enzymology
        (Biochemistry and Molecular Biophysics); Nutrition; Urinary System
        (Chemical Coordination and Homeostasis)
ΙT
     Parts, Structures, & Systems of Organisms
        kidney: excretory system; nephron: excretory system
ΙT
     Diseases
        hypertension: vascular disease
TΤ
     Chemicals & Biochemicals
        calcium: dietary intake; potassium-potassium-ATPase: activity, alpha-1
        isoform, membrane ion translocating pump, renal, gene
        expression 1
ΙT
     Methods & Equipment
        RT-PCR [reverse transcriptase-polymerase chain reaction]:
        analytical method
ORGN Super Taxa
        Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        rat (Muridae): strain-Wistar-Kyoto, strain-spontaneously hypertensive,
        weanling
```

39.

ORGN Organism Superterms

Animals; Chordates; Mammals; Nonhuman Mammals; Nonhuman Vertebrates; Rodents; Vertebrates

RN 9000-83-3 (ATPASE)

L15 ANSWER 17 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

AN 1998:29918 BIOSIS

DN PREV199800029918

- TI Regulation of phagosomal acidification. Differential targeting of Na+/H+ exchangers, Na+/K+-ATPases, and vacuolar-type H+-ATPases.
- AU Hackam, David J.; Rotstein, Ori D.; Zhang, Wei-Jian; Demaurex, Nicolas; Woodside, Michael; Tsai, Olivia; Grinstein, Sergio (1)
- CS (1) Division Cell Biol., Hosp. Sick Children, 555 University Ave., Toronto, ON M5G 1X8 Canada
- SO Journal of Biological Chemistry, (Nov. 21, 1997) Vol. 272, No. 47, pp. 29810-29820. ISSN: 0021-9258

DT Article

LA English

Vacuolar-type (V) ATPases are thought to be the main determinant of AB phagosomal adidification. In phagosomes containing mycobacteria, which ostensibly impair the delivery of V-ATPases to the phagosomal membrane, the pH would be expected to be near neutral. This prediction was tested by microfluorescende ratio imaging using macrophages from mice susceptible to mycobacterial infection. Although less acidic than their counterparts containing dead bacteria, phagosomes containing live Mycobacteria bovis were nearly long unit more acidic than the cytosol, suggesting the existence of alternate H+ transport mechanisms. We therefore investigated whether Na+/H+ exchange (NHE) contributes to phagosomal acidification: Immunoblotting, reverse transcriptase-polymerase chain reaction and pharmacological studies indicated that NHE1 is the predominant isoform of the exchanger in macrophages. Fractionation revealed that NHE1 is incorporated into the phagosomal membrane, and measurements of pH indicated that it is functional in this location. Nevertheless, acidification of the lumen of phagosomes containing either latex beads or live M. bovis was insensitive to (3-methylsulfonyl-4piperidinoben zoyl)-guanidine methanesulfonate, a potent inhibitor of NHE 1. This may have been due to the absence of an appropriate lumen to cytosol Na+ ďřådient, because the phagosomal membrane was found to be devoid of Native pumps. Unexpectedly, the acidification of M. bovis phagosomes was fully reversed by specific inhibitors of the vacuolar H+ATPase, suggesting that ATPases are present only transiently or in reduced quantities in the phagosomal membrane.

Alternatively, lacid equivalents accumulated in endosomes by V-ATPases may be delivered to the mycobacterial phagosome by carrier vesicles devoid of ATPases.

IT Major Concepts

Enzymology (Biochemistry and Molecular Biophysics); Infection; Membranes (Cell Biology)

IT Parts, Structures, & Systems of Organisms

macrophage immune system; phagosome: acidification, membrane

IT Diseases

Mycobacterium-bovis infection: bacterial disease

IT Chemicals & Biochemicals

133

sodium/hydrogen exchanger; sodium/potassium ATPase; vacuolar-type hydrogen ATPase

ORGN Super Taxa

Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia; Mycobacteriaceae: Mycobacteria, Actinomycetes and Related Organisms, Eubacteria, Bacteria, Microorganisms ORGN Organism Name

mouse (Muridae): host; J774 (Muridae); Mycobacterium-bovis (Mycobacteriaceae): pathogen

ORGN Organism Superterms

Animals; Baggeria; Chordates; Eubacteria; Mammals; Microorganisms; Nonhuman Mammals; Nonhuman Vertebrates; Rodents; Vertebrates

RN 9000-83-3 (ATPASE)

- ANSWER 18 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15
- 1997:482937 BTOSIS ΑN
- PREV199799782140 DN
- The apoptosis mecrosis paradox: Apoptogenic proteases activated after ΤI mitochondrial permeability transition determine the mode of cell death.
- Hirsch, Tamara, Marchetti, Philippe; Susin, Santos A.; Dallaporta, Bruno; ΑU Zamzami, Naoutal; Marzo, Isabel; Geuskens, Maurice; Kroemer, Guido
- Centre Natl. Rech. Sci., Unite Propre Recherche 420, 19 rue Guy Moquet, F-94801 Villejuif France CS
- Oncogene, (1997) Vol. 15, No. 13, pp. 1573-1581. ISSN: 0950-9232 SO
- DT Article
- LA English
- Mitochondrial alterations including permeability transition (PT) AΒ constitute critical events of the apoptotic cascade and are under the control of Bc1-2 related gene products. Here we show that induction of PT is sufficient to activate CPP32-like proteases with DEVDase activity and the associated cleavage of the nuclear DEVDase substrate poly(ADP-ribose) polymerase (PARP). Thus, direct intervention on mitochondria using a ligand of the mitochondrial benzodiazepin receptor or a protonophore causes DEVDase activation. In addition, the DEVDase activation triggered by conventional apoptosis inducers (glucocorticoids or topoisomerase inhibitors) is prevented by inhibitors of PT. The protease inhibitor N-benzyloxycabonyl-Val-Ala-Asp-fluoromethylketone (Z-VAD.fmk) completely prevents the activation of DEVDase and PARP cleavage, as well as the manifestation of nuclear apoptosis (chromatin condensation, DNA fragmentation, hypoploidy). In addition, Z-VAD.fmk delays the manifestation of apoptosis-associated changes in cellular redox potentials (hypergeneration of superoxide anion, oxidation of compounds of the inner mitochondrial membrane, depletion of non-oxidized glutathione), as well as the exposure of phosphatidylserine residues in the outer plasma membrane leaflet. Althourgh z-VAD.fmk retards cytolysis, it is incapable of preventing disruption of the plasma membrane during protracted cell culture (12-24 h), even in conditions in which it completely blocks nuclear apoptosis (chromatin condensation and DNA fragmentation) Electron microscopic analysis confirms that cells treated with PT inducers alone undergo apoptosis, whereas cells kept in identical conditions in the presence of Z-VAD.fmk die from necrosis. These observations are compatible with the hypothesis that PT would be a rate limiting step in both the apoptotic and the necrotic modes of cell death. In contrast, it would be the availability of apoptogenic proteases that would determine the choice between the two death modalities.

ITMajor Concepts

> Blood and Lymphatics (Transport and Circulation); Endocrine System (Chemical Coordination and Homeostasis); Enzymology (Biochemistry and Molecular Biophysics); Genetics; Membranes (Cell Biology); Metabolism

- ΙT Chemicals & Biochemicals
- PROTEASES; PROTEASE
  Miscellaneous Descriptors IT

ANIMAL MODEL APOPTOGENIC PROTEASE ACTIVATION; APOPTOSIS-NECROSIS PARADOX; ÇELL BIOLOGY; CELL DEATH MODE; ENDOCRINE SYSTEM; IMMUNE

SYSTEM; IN TRO MODEL SYSTEM; MITOCHONDRIAL PERMEABILITY TRANSITION; MOLECULAR GENETICS; THYMOCYTES

ORGN Super Taxa

Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

Balb/C mouse (Muridae)

ORGN Organism Superterms

animals; chordates; mammals; nonhuman mammals; nonhuman vertebrates; rodents; vertebrates

RN 9014-01-1 (PROTEASES) 9001-92-7 (PROTEASE)

- L15 ANSWER 19 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1997:463912 BIÓSIS
- DN PREV199799763115
- TI Synthetically modified cellulose: An alternative to synthetic membranes for wise in haemodialysis.
- AU Hoenich, Nicholas A. (1); Woffindin, Celia; Stamp, Susan; Roberts, Sarah J.; Turnbull, Jean
- CS (1) Dep. Med. Sch. Clin. Med. Sci., Floor 4, William Leech Build., Med. Sch., Univ. Newcastle, Framlington Place, Newcastle upon Tyne NE2 4HH UK
- SO Biomaterials, (1997) Vol. 18, No. 19, pp. 1299-1303. ISSN: 0142-9612.
- DT Article
- LA English
- AB Renal replacement therapy relies predominantly on the use of cellulose-based membranes. Such membranes have a biocompatibility profile which is infector to membranes manufactured from synthetic polymers. Synthetically modified cellulose (SMC) is a new, low-flux haemodialysis membrane in which hydroxyl groups have been replaced with benzyl groups. The biocompatibility profile characterized by changes in white cell and platelet counts and the activation of complement components (C3a, C5a and C5b-9) have been studied in vivo and compared with those of cellulose acetate, unmodified cellulose (Cuprophan) and low-flux polysulphone (Fresenius Polysulfone) in the same group of patients. For SMC, the white cell count at 15 min declined to 65.6% of pretreatment level, compared with 63.8% for the cellulose acetate, 79.6% for low-flux polysulphone and 28.1% for Cuprophan, thereafter returning to pretreatment levels. Both modified cellulose membranes were superior to unmodified cellulose (P = 0.001); the differences between the modified cellulose membranes were not significant statistically. The changes induced by allithree cellulose-based membranes exceeded those for low-flux polysulphone (P=0.001). Associated with the neutropenia was a reduction in platelet count, but this was independent of membrane type. The mean time-averaged concentrations of C3-des Arg over 150 min were 1168 ng ml-1 (SMC), 1030  $n\ddot{g}$  ml-1 (cellulose acetate), 1297 ng ml-1 (Cuprophan) and 790 ng ml-1 (low-flux polysulphone). Equivalent values for C5-des Arg were 6.12 (SMC), 2.98 (cellulose acetate), 11.03 (Cuprophan) and 1.33 ng ml-1 (low-flux polysulphone). C5b-9 values were 385 (SMC), 386 (cellulose acetate), 177% (Cuprophan) and 185 ng ml-1 (low-flux polysulphone). For each of the complement components the differences between the membranes were significant (P = 0.0009 (C3a-des Arg), P = 0.0001 (c5a-des Arg and C5b-9)). The levels of C5b-9 generated during dialysis also showed a significant positive correlation compared to C5a for all membranes considered as a single group (Pearson's correlation coefficient = 0.870, P = 0.0001). It is concluded that the modification of the cellobiosic unit is a promising approach to improve the biocompatibility profile of cellulose-based membranes. The two different methods of modification lead to similar improvements in biocompatibility compared with unmodified cellulose, but as yet do not match that of low-flux polysulphone.

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ΙT
     Major Concepts
        Biochemistry and Molecular Biophysics; Blood and Lymphatics (
        Transport and Circulation); General Life Studies; Immune System
        (Chemical Coordination and Homeostasis); Methods and Techniques;
        Pathology: $\psi rinary System (Chemical Coordination and Homeostasis)
     Chemicals & Biochemicals
ΙT
        CELLULOSE .
     Miscellaneous Descriptors
ΙT
        BIOBUSINESS; BIOCHEMISTRY AND BIOPHYSICS; BIOMATERIALS; CELLULOSE;
        COMPLEMENT; COMPLEMENT ACTIVATION; HEMODIALYSIS; MEDICAL RESEARCH;
        METHODOLOGY, PATIENT; RENAL REPLACEMENT THERAPY; SYNTHETIC
        MEMBRANE ALTERNATIVES; SYNTHETICALLY MODIFIED;
        THERAPEUTIC
ORGN Super Taxa
        oer Taxa (1987)
Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Hominidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
     9004-34-6 (CELLULOSE)
RN
     ANSWER 20 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
     1997:435991 BİOSIS
AN
DN
     PREV199799735194
     The major intrinsic protein family of arabidopsis has 23 members that form
ΤI
     three distinct groups with functional aquaporins in each group.
     Weig, Alfons; Deswarte, Corine; Chrispeels, Maarten J. (1) (1) Dep. Biol 20116, University Calif. San Diego, 9500 Gilman Dr., La
ΑU
CS
     Jolla, CA 92093-0116 USA
     Plant Physiology (Rockville), (1997) Vol. 114, No. 4, pp. 1347-1357.
SO
     ISSN: 0032-0889
DT
     Article
LA
     English
     Aquaporins, proteins that enhance the permeability of
AΒ
     biological membranes to water, are widely distributed in living
     organisms. They are 26- to 29-kD proteins that belong to the major
     intrinsic protein (MIP) family of channels. By searching the Arabidopsis
     thaliana expressed sequence tag database and by using the
     polymerase chain reaction with oligonucleotides to conserved plant
     aquaporin domains, we identified 23 expressed Arabidopsis MIP genes. Eight
     of these had been previously identified as active aquaporins, and two
     additional ones are now reported to have water-transport
     activity in Xenopus laevis oocytes. One of these is highly expressed in
     suspension-cultured cells. On a dendrogram these 23 MIP sequences cluster
     into three groups: the first group has 11 members and contains the plasma
     membrane aquaporins, the second group also has 11 members and contains the
     tonoplast aquaporins, and the third group has only a single member. This
     MIP protein, provisionally called At-NLM1, is most closely related to the
     Gm-NOD26 protein that is found in the bacteroid membranes of soybean
     (Glycine max (L)) nodules; At-NLM1 is an active aquaporin when expressed in
     oocytes. With a semiquantitative slot-blot analysis technique, we
     determined the expression levels of 22 MIP genes in the various organs.
     The quantitative polymerase chain reaction was used to determine
     the effects of \frac{1}{2} various stress treatments on the expression of NLM1.
     Major Concepts

Biochemistry and Molecular Biophysics; Cell Biology; Genetics;

Membranes (Cell Biology)
TT
IT
     Miscellaneous Descriptors
        AQUAPORINS; BIOCHEMISTRY AND BIOPHYSICS; GENETIC METHOD; MAJOR
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INTRINSIC PROTEIN GENES; MEMBRANE PROTEINS; MEMBRANES

; MIP GENES; MOLECULAR GENETICS; QUANTITATIVE POLYMERASE CHAIN REACTION; WATER-SELECTIVE CHANNELS

ORGN Super Taxa

Cruciferae Dicotyledones, Angiospermae, Spermatophyta, Plantae; Leguminosae Dicotyledones, Angiospermae, Spermatophyta, Plantae ORGN Organism Name

Arabidopsis thaliana (Cruciferae); Glycine max (Leguminosae) ORGN Organism Superterms

angiosperms; dicots; plants; spermatophytes; vascular plants

- ANSWER 21 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15
- ΑN 1997:409938 BÍOSIS
- DN PREV199799701981
- G-protein-mediated signaling in cholesterol-enriched arterial smooth ΤI muscle cells. 1. Reduced membrane-associated G-protein content due to diminished isoprenylation of G-gamma subunits and p21ras.
- ΑU Pomerantz, Kenneth B.; Lander, Harry M.; Summers, Barbara; Robishaw, Janet
- D.; Balcueva, Eric; Hajjar, David P. (1) (1) Dep. Med. Biochem., Cornell Univ. Med. Coll., 1300 York Ave., New CS York, NY 10021 USA
- Biochemistry (1997) Vol. 36, No. 31, pp. 9523-9531. SO ISSN: 0006-2960
- DT Article
- LAEnglish
- Mechanisms contributing to altered heterotrimeric G-protein expression and AB subsequent signaling events during cholesterol accretion have been unexplored. The influence of cholesterol enrichment on G-protein expression was examined in cultured smooth muscle cells that resemble human atheros@lerotic cells by exposure to cationized LDL (cLDL). cLDL, which increases cellular free and esterified cholesterol 2-fold and 10-fold, respectively, reduced the cell membrane content of G-alpha-i-1, G-alpha-i-2, Gralpha-i-3, Gq/11, and G-alpha-s. The following evidence supports the premise that the mechanism by which this occurs is due to reduced isoprehýlation of the G-gamma-subunit. First, the inhibitory effect of cholesterol enrichment on the membrane content of G-alpha-i subunits was found to be post-transcriptional, since the mRNA steady-state levels of G-alpha-i(1-3) were unchanged following cholesterol enrichment. Second, the membrane expression of alpha and beta subunits was mimicked by cholesterol and 17-ketocholesterol, both of which inhibit HMG-CoA reductase. Third, inhibition of G-alpha-i and G-beta expression in cholesterol-enriched cells was overcome by mevalonate, the immediate product of HMG COA reductase. Fourth, pulse-chase experiments revealed that cholesterol enrichment did not reduce the degradation rate of membrane-associated G-alpha-i subunits. Fifth, cholesterol enrichment also reduced membrahe expression of G-gamma-5, G-gamma-7-upper; these gamma subunits are résponsible for trafficking of the heterotrimeric G-protein complex to the cell membrane as a result of HMG-CoA reductase-dependent post-translat မြို့စိုက်ချ lipid modification (geranylgeranylation) and subsequent membrane association. Cholesterol enrichment did not alter expression of G-gamma-5 mRNA as assessed by reverse transcriptase polymerase chain reaction supporting a post-transcriptional defect in G-gamma subunit expression. Fifth, cholesterol enrichment also reduced the membrane content of p21ras (a low molecular weight G-protein requiring farnesylation for membrane targeting) but did not alter the membrane content of the two proteins that do not require isoprenylation for membrane association-PDGF-receptor or p60-src. Reduced G-protein confent in cholesterol-laden cells was reflected by reduced G-protein-mediated signaling events, including ATP-induced GTPase activity, thrombin-induced inhibition of cyclic AMP accumulation, and MAP kinase activity. Collectively, these results demonstrate that cholesterol

enrichment reduces G-protein expression and signaling by inhibiting isoprenylation and subsequent membrane targeting. These results provide a molecular basis for altered G-protein-mediated cell signaling processes in cholesterol-enriched cells. Major Concepts Biochemistry and Molecular Biophysics; Cardiovascular System ( IT Transport and Circulation); Cell Biology ΙT Chemicals & Biochemicals CHOLESTEROL; CYCLIC AMP Miscellaneous Descriptors IT ARTERIAL SMOOTH MUSCLE CELLS; BIOCHEMISTRY AND BIOPHYSICS; CARDIOVASCÜLAR SYSTEM; CELL SIGNALING; CHOLESTEROL; CIRCULATORY SYSTEM; CYCLIC AMP; EXPRESSION; G-GAMMA-5 MESSENGER RNA; G-GAMMA-5 MRNA; G-PROTEIN; LOW DENSITY LIPOPROTEIN; MEMBRANE ASSOCIATED; MEMBRANES; P21RAS ORGN Super Taxa Leporidae Lagomorpha, Mammalia, Vertebrata, Chordata, Animalia; Muridae: Rodentia, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name rabbit (Leporidae); rat (Muridae) ORGN Organism Superterms animals; chordates; lagomorphs; mammals; nonhuman mammals; nonhuman vertebrates; rodents; vertebrates 57-88-5 (CHOLESTEROL) RN 60-92-4 (CYCLIC AMP) ANSWER 22 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15 1996:260673 BIOSIS ΑN PREV199698816802 DN ΤI Alterations in adhesion, transport, and membrane polymers in adhesion-deficient pseudomonads. Oppenheimer, Stephanie R. (1); Condee, Charles C.; Fletcher, Madilyn (1); ΑIJ Deflaun, Mary F. CS (1) Cent. Marine Biotechnol., Univ. Maryland Biotechnol. Inst., Baltimore, MD 21202 USA Abstracts of the General Meeting of the American Society for Microbiology, SO (1996) Vol. 96, No. 0, pp. 405. Meeting Info : 96th General Meeting of the American Society for Microbiology New Orleans, Louisiana, USA May 19-23, 1996 ISSN: 1060-2011. DT Conference LA English IT Major Concepts! Biochemistry and Molecular Biophysics; Cell Biology; Membranes (Cell Biology); Methods and Techniques; Physiology Miscellaneous Descriptors IT ANALYTICAE METHOD; CELL SURFACE CHARACTERISTICS; ELECTROPHORESIS; LATERAL DISPERSION; MEETING ABSTRACT; OUTER MEMBRANE PROTEINS; WILD TYPE ORGN Super Taxa Pseudomonadaceae: Eubacteria, Bacteria ORGN Organism Name Pseudomonadaceae (Pseudomonadaceae) ORGN Organism Superterms bacteria; eubacteria; microorganisms ANSWER 23 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15 AN 1996:123662 BIOSIS DN PREV199698695797 TΙ Laboratory markers as an index of aging.

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ΑU
     Narayanan, Sheshadri N.
     Dep. Pathol. New York Med. Coll., Metropol. Hosp. Cent., New York, NY
CS
     10029 USA
     Annals of Clinical and Laboratory Science, (1996) Vol. 26, No. 1, pp.
SO
     ISSN: 0091-7370.
DT
     Article
LA
     English
     At the cellular level, mutations in deoxyribonucleic acid (DNA) can lead
AB
     to synthesis of altered proteins which are unable to sustain specific cell
     functions, eventually leading to its death. Veritably apoptosis, or
     programmed cell death, is a device to eliminate heavily mutated cells.
     Cell membranes with altered proteins can be recognized
     as foreign by the immune system, thus triggering autoimmunity. Molecular
     biology techniques allow us to examine changes that occur in DNA,
     reflected by polymorphisms and variable numbers of tandem repeats (VNTR).
     A general decline in organ function is associated with aging. However,
     these changes may also be precipitated by disease processes. Homeostatic
     control by the hypothalamus-pituitary-adrenal axis is also compromised
     with aging, leading to an increase in plasma adrenocorticotrophic hormone
     (ACTH) and cofficosteroid levels. Derangement of the immune system with
     aging results in dysregulation of cytokine production. The ability of the
     cell to survive the onslaught of oxygen-free radicals with enzymatic and
     nonenzymatic antioxidants, and to repair DNA by activation of nuclear
     enzymes such as poly (ADP-ribose) polymerase (PAD-PRP), are some
     of determinanț of aging.
ΙT
     Major Concepts
        Biochemistry and Molecular Biophysics; Blood and Lymphatics (
        Transport and Circulation); Cell Biology; Clinical Chemistry
        (Allied Medical Sciences); Clinical Immunology (Human Medicine, Medical
        Sciences) Endocrine System (Chemical Coordination and Homeostasis); Enzymology (Biochemistry and Molecular Biophysics); Geriatrics (Human
        Medicine, Medical Sciences); Membranes (Cell Biology);
        Metabolism Pathology; Toxicology
     Chemicals & Biochemicals
IT
        OXYGEN-FREE RADICAL; POLY (ADP-RIBOSE) POLYMERASE; ACTH
ΙT
     Miscellaneous Descriptors
        ANTIOXIDANT; APOPTOSIS; AUTOIMMUNITY; CELL MEMBRANE; CYTOKINE
        PRODUCTION; DNA MUTATION; ENZYMATIC ANTIOXIDANT; NUCLEAR ENZYME
        ACTIVATION; OXYGEN-FREE RADICAL; PLASMA ACTH LEVEL; POLY (ADP-RIBOSE)
        POLYMERASE; SERUM CORTICOSTEROID LEVEL
ORGN Super Taxa
        Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia
ORGN Organism Name
        human (Homimidae)
ORGN Organism Superterms
        animals; chordates; humans; mammals; primates; vertebrates
     11062-77-4 (OXYGEN-FREE RADICAL)
RN
     9055-67-8 (POLY (ADP-RIBOSE) POLYMERASE)
     9002-60-2 (ACTE)
     ANSWER 24 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
L15
ΑN
     1996:31359 BIOSIS
DN
     PREV199698603494
ΤI
     Deoxygenation induced alterations in sickle cell
     membrane cholesterol exchange.
     Kavecansky, Jüřáj; Schroeder, Friedhelm; Joiner, Clinton H. (1)
ΑU
CS
     (1) Children's Hosp. Med. Center, Comprehensive Sickle Cell Center, 3333
     Burnet Ave., Cincinnati, OH 45229-3039 USA
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American Journal of Physiology, (1995) Vol. 269, No. 5 PART 1, pp.

SO

C1105-C1111. ISSN: 0002-9513

DT Article

LA English

Changes in a membrane sterol exchange of sickle red blood cells (SS RBC) AΒ induced by deokygenation were studied using the fluorescent cholesterol analogue dehydroergosterol (DHE), DHE uptake by SS RBC membrane was measured by the incubation of SS RBC with small unilamellar vesicles (SUV) containing DHE Deoxygenation of SS RBC, but not normal RBC, increased the rate of DHE uptake. DHE membrane content after 5 h of incubation with SUV in the cell-to SUV ratio of 1:1 (mol lipid) was 16.25 +- 0.94 and 12.22 +- 0.85% of total sterol for deoxygenated and oxygenated cells, respectively. Membrane spicules isolated from these deoxygenated SS RBC had threefold higher DHE content, suggesting that the increased sterol exchange was localized to spicules. When isolated spicules were incubated with DHE-SUV directly, 91 44 3% of membrane sterol was rapidly exchanged, in contrast to intact RBC, in which a maximum of 33% of sterol could be exchanged. The results suggest that spicule formation in SS RBC alters membrane cholesterol structure, such that a domain of cholesterol that is normally nonexchangeable becomes readily exchangeable with exogenous sterol.

Major Concepts IT

> Blood and Lymphatics (Transport and Circulation); Cell Biology; Membranes (Cell Biology); Metabolism; Morphology

Chemicals & Biochemicals IT

CHOLESTEROL

IT

Miscellaneous Descriptors HEMOGLOBIN S POLYMERIZATION; UNILAMELLAR VESICLE

ORGN Super Taxa

Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

human (Hominidae)

ORGN Organism Superterms

animals; chordates; humans; mammals; primates; vertebrates

57-88-5 (CHOLESTEROL) RN

- ANSWER 25 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15
- 1995:402152 BIOSIS AN
- DN PREV199598416452
- Evaluation by histology, immunohistology and PCR of protocollized renal ΤI biopsies 1 week post-transplant in relation to subsequent rejection
- Kooijmans-Coutinho, M. F. (1); Bruijn, J. A.; Hermans, J.; Schindler, R.; AU Frei, U.; Schrama, E.; Van Es, L. A.; Daha, M.r.; Van Der Woude, F. J.
- (1) Leiden Un v Hosp., Dep. Nephrol., Build. 1, C3-P, P.O. Box 9600, 2300 CS RC Leiden Netherlands
- Nephrology Dialysis Transplantation, (1995) Vol. 10, No. 6, pp. 847-854. SO ISSN: 0931-0509.
- DT Article
- LAEnglish
- Renal biopsies were performed 1 week following renal transplantation at a AR time without clinical evidence of rejection in 43 patients (13 females, mean age 48 years range 18-60 and 30 males, mean age 43 years range 17-59 years). Thirty is ix biopsies were available for histological or immunohistochemical analysis. Immunohistochemical analyses were performed with monoclonal antibodies against leukocytes (CD45), monocytes (WT14), complement factor 3 (C3), T-cells (Leu4), T-cell receptor alpha-beta and gamma-delta, tumour necrosis factor alpha (TNF-alpha), IL-2 receptor (IL2-R, TAC), intercellular adhesion molecule-1 (ICAM1) and HLA-DR. The slides were scored semiquantitatively with the observers having no

knowledge of clinical or patient data. TNF-alpha and IL-2R were also measured by quantitative PCR. None of the studied parameters correlated to delayed graft function or graft loss. Histological analysis showed that both focal interstitial infiltrate (18/35) and tubular basement membrane disruption (11/35) were followed by a higher incidence of subsequent rejection (P = 0.03 and 0.02 respectively). Also positivity for WT14 around tubuli (P=0.02) was associated with subsequent occurrence of rejection. The intensity of staining of ICAM-1 on PTC as well as TAC on proximal tubular cells was associated with the number of subsequent rejection episodes. The association between the IL-2 receptor and subsequent rejection was also found applying PCR to the tissue specimens. We conclude that the presence of focal interstitial infiltrates and tubulitis in 1-week biopsies from well-functioning grafts carries an increased risk of subsequent rejection. The observed infiltrate outside the tubuli may consist of either monocytes or lymphocytes. Further studies, both in vitro and in vivo, applying immunohistochemical and molecular biological techniques will be necessary to further elucidate the role of adhesion molecules and interleukins in early and ongoing rejection.

IT Major Concepts

Blood and Lymphatics (Transport and Circulation); Cell Biology; Clinical Immunology (Human Medicine, Medical Sciences); Endocrine System (Chemical Coordination and Homeostasis); Enzymology (Biochemistry and Molecular Biophysics); Immune System (Chemical Coordination and Homeostasis); Membranes (Cell Biology); Methods and Techniques; Physiology; Surgery (Medical Sciences); Urology (Human Medicine, Medical Sciences)

IT Miscellaneous Descriptors

INTERLEUKÉN 2 RECEPTOR; INTRACELLULAR ADHESION MOLECULE-I; LYMPHOCYTE; MACROPHAGE; MONOCYTE; POLYMERASE CHAIN REACTION; RENAL TRANSPLANTATION

ORGN Super Taxa

Hominidae Primates, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

human (Hominidae)

ORGN Organism Superterms

animals; chordates; humans; mammals; primates; vertebrates

- L15 ANSWER 26 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1994:315130 BIOSIS
- DN PREV199497328130
- TI Twenty seven nucleotide deletion within exon 11 of the erythrocyte band 3 gene in Indonesian ovalocytosis.
- AU Takeshima, Yasuhiro (1); Sofro, Abdul Salam; Suryantoro, Purnomo; Narita, Naoko (1); Matsuo, Masafumi (1)
- CS (1) Div. Genetics, International Center Med. Res., Kobe University Sch. Med., Kobe Japan
- SO Japanese Journal of Human Genetics, (1994) Vol. 39, No. 1, pp. 181-185. ISSN: 0021-50743
- DT Article
- LA English
- AB We here report the molecular characterization of an Indonesian ovalocytosis. The analysis of genomic gene by polymerase chain reaction shows that the individual has two amplified products from a region encompassing exon 11 of the erythrocyte band 3 gene. The sequence of the larger product matched completely with that of normal individuals. In the sequence of the smaller product, 27 nucleotides within exon 11 disappeared. The deletion removes a total of nine amino acids in the boundary of cytoplasmic and membrane domains of band 3 protein, a membrane anion transporter protein. This is the first report to confirm

#### 09/755,701 Tran

the heterogeneous presence of an altered membrane band 3 protein in indonesian ovalocytosis.

ΙT Major Concepts

Anthropology, Blood and Lymphatics (Transport and Circulation; Cell Biology; Enzymology (Biochemistry and Molecular Biophysics); Genetics; Membranes (Cell Biology)

ΙT Miscellaneous Descriptors

> MEMBRANE ANION TRANSPORTER PROTEIN; MOLECULAR CHARACTERIZATION; POLYMERASE CHAIN REACTION

ORGN Super Taxa

Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

human (Hominidae)

ORGN Organism Superterms

animals; chordates; humans; mammals; primates; vertebrates

- L15 ANSWER 27 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.
- AN 1992:231281 BIOSIS
- DN BA93:119306
- DEFECTIVE ANION TRANSPORT ACTIVITY OF THE ABNORMAL BAND 3 IN ΤI HEREDITARY OVALOCYTIC RED BLOOD CELLS.
- ΑU
- CS
- SCHOFIELD A EMREARDON D M; TANNER M J A DEP. BIOCHEM. SCH. MED. SCI., UNIV. BRISTOL, BRISTOL BS8 1TD, UK. NATURE (LOND); (1992) 355 (6363), 836-838. CODEN: NATURES ISSN: 0028-0836. SO
- FS BA; OLD
- LA English
- Hereditary ovalocytosis is common in some areas of Melanesia and South AB East Asia where malaria is endemic. These red cells resist invasion by malarial parasites in vitrol, 2 and ovalocytic individuals are less parasitized than normal3. This has been attributed to greater rigidity of ovalocytic red cells4,5. It has been suggested that South East Asia ovalocytosis results from the heterozygous presence of an altered membrane anion transporter (band 3)6,7. We have used the polymerase chain reaction to clone the abnormal band 3 complementary DNA from an ovalocytic of Indian origin8 and found two changes from the normal protein: a point mutation (Lys 56 .fwdarw. Glu) and the deletion of the sequence AFSPQVLAA (residues 400-408), but no evidence for an N-terminal extension7. The deletion is also found in the abnormal band of South East Asian ovalocytes9 and seems to be responsible for the unusual properties of the ovalocytic red cell. We show here that the membrane domain of the abnormal ovalocyte band 3 has a substantially altered structure and that the protein is defective in anion transport activity. The changed transport properties of the red cells may have a role in the reduced parasitaemia of ovalocytic individuals.
- Miscellaneous Descriptors TΨ

HUMAN ALTERED MEMBRANE FUNCTION POINT MUTATION SEQUENCE DÉLÉTION ENDEMIC MALARIA REGION REDUCED PARASITEMIA POLYMERASE CHAIN REACTION MELANESIA SOUTHEAST ASIA

- ANSWER 28 OF 29 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. L15
- AN 1989:221221 BIOSIS
- DN BA87:112838
- SUPRAMOLECULAR SURFACTANTS AMPHIPHILIC POLYMERS DESIGNED TO ΤI DISRUPT LIPID MEMBRANES.
- ΑU REGEN S L; JAYASURIYA N; FABIANOWSKI W

A Company of the Comp

- CS DEP. CHEM., LEHIGH UNIV., BETHLEHEM, PA. 18015.
- BIOCHEM BIOPHYS RES COMMUN, (1989) 159 (2), 566-571. CODEN: BBRCA9 ISSN: 0006-291X.

FS BA; OLD

English

LA

AB Simple polyesters derived from poly(ethylene glycol)s and .alpha., .omega.-dicarboxylic acids exhibit a broad range of activity in disrupting phospholipid membranes. This activity has been analyzed by measuring the release of liposome-encapsulated 5(6)-carboxy-filiorescein (CF). Comparison with an analogous monomeric surfactant, and with Triton X-100, demonstrate that macromolecular activity is a sensitive function of the size of the hydrophobic and hydrophilic segments within each repeat unit, and that high disrupting power is possible. In vitro studies with the human immunodeficiency virus type-1 have revealed that those polyesters which exhibit the highest membrane disrupting power also provide significant protection for human CD4+ lymphocytes against HIV-1. The potential for adjusting and utilizing these "supramolecular surfactants" in medicine is briefly discussed.

IT Miscellaneous Descriptors
HUMAN IMMUNODEFICIENCY VIRUS LYMPHOCYTE VIRAL DEACTIVATION

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AN 1980:165122 BÌOSIS

DN BA69:40118

TI PHOTO REGULATION OF THE INCORPORATION OF GUAIACYL UNITS INTO LIGNINS.

AU GRAND C; RANJEŸA R; BOUDET A M; ALIBERT G

CS CENT. PHYSION VEG., LAB. ASSOC. FRANCE, CNRS 241, LL8 ROUTE NARBONNE, F-3L077 TOULOUSE CEDEX, FR.

SO PLANTA (BERL); (1979) 146 (3), 281-286. CODEN: PLANAB ISSN: 0032-0935.

FS BA; OLD

LA English

When fed with [14C] phenylalanine in the light, xylem tissues isolated from poplar (Populus .times. euramericana) stems were able to incorporate part of the radioactivity into both the guaiacyl and the syringyl residues of lignins. In the dark, only syringyl units were integrated into the polymer whereas the guaiacyl residues remained unlabeled. When a membrane perturber (isopropanol) was added to the incubation mixture, the label was incorporated into the guaiacyl units either in the light or in the dark. Conversely, a membrane stabilizer (CaCl2) prevented the labeling of the guaiacyl units even when the tissues were illuminated. These results suggest that light acts through the modification of membrane permeabilities, altering specifically the synthesis and the transport or the polymerization of guaiacyl-type units during the process of lignification.

IT Miscellaneous Descriptors

POPULUS-EURAMERICANA CARBON-14 PHENYL ALANINE **MEMBRANE** PERMEABILIŢŸ SYNTHESIS **TRANSPORT** 

RN 9005-53-2D (LIĞNINS) 14762-75-5 (ÇARBON-14) 63-91-2Q, 3617-44-5Q (PHENYL ALANINE)